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(\$4) Liffe: HUMAN ANTIBODIES THAT BIND HUMAN IL-12 AND METHODS FOR PRODUCING

(57) Abstract

recombinant human antibodies, are also encompassed by the invention. Mucleic acids, vectors and host cells for expressing the recombinant human antibodies of the invention, and methods of synthesizing the hIL-12 and for inhibiting hIL-12 activity, e.g., in a human subject suffering from a disorder in which hIL-12 activity is detrimental. a full-length antibody or an antigen-binding portion thereof. The antibodies, or antibody portions, of the invention are useful for detecting Preferred antibodies have high affinity for hIL-12 and neutralize hIL-12 activity in vitro and in vivo. An antibody of the invention can be Human antibodies, preferably recombinant human antibodies, that specifically bind to human interleukin-12 (htL-12) are disclosed.

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WAN ANTIBODIES THAT BIND HUMAN IL-12

Related Applications

This application is a non-provisional application claiming priority to U.S. provisional application Serial No. 60/126,603, filed March 25, 1999, the contents of which are hereby incorporated by reference.

Background of the Invention

Human interleukin 12 (IL-12) has recently been characterized as a cytokine with a unique structure and pleiotropic effects (Kobayashi, et al. (1989) J. Exp Med. 170:827-845; Seder, et al. (1993) Proc. Natl. Acad. Sci. 90:10188-10192; Ling, et al. (1995) J. Exp Med. 154:116-127; Podlaski, et al. (1992) Arch. Biochem. Biophys. 294:230-237).

IL-12 plays a critical role in the pathology associated with several diseases involving immune and inflammatory responses. A review of IL-12, its biological activities, and its role in disease can be found in Gately et al. (1998) Ann. Rev. Immunol. 16: 495-521.

Structurally, IL-12 is a heterodimeric protein comprising a 35 kDa subunit (p35)

and a 40 kDa subunit (p40) which are both linked together by a disulfide bridge (referred to as the "p70 subunit"). The heterodimeric protein is produced primarily by antigenpresenting cells such as monocytes, macrophages and dendritic cells. These cell types also secrete an excess of the p40 subunit relative to p70 subunit. The p40 and p35 subunits are genetically unrelated and neither has been reported to possess biological activity, although the p40 homodimer may function as an IL-12 antagonist.

Functionally, IL-12 plays a central role in regulating the balance between antigen

specific T helper type (Th1) and type 2 (Th2) lymphocytes. The Th1 and Th2 cells govern the initiation and progression of autoimmune disorders, and IL-12 is critical in the regulation of Th₁-lymphocyte differentiation and maturation. Cytokines released by the Th1 cells are inflammatory and include interferon y (IFNy), IL-2 and lymphotoxin (LT). Th2 cells secrete IL-4, IL-5, IL-6, IL-10 and IL-13 to facilitate humoral immunity, allergic reactions, and immunosuppression.

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Consistent with the preponderance of Th1 responses in autoimmune diseases and the proinflammatory activities of IFN γ , IL-12 may play a major role in the pathology associated with many autoimmune and inflammatory diseases such as rheumatoid arthritis (RA), multiple sclerosis (MS), and Crohn's disease.

documented by p40 mRNA levels in acute MS plaques. (Windhagen et al., (1995) J. Exp. Med. 182: 1985-1996). In addition, ex vivo stimulation of antigen-presenting cells with CD40L-expressing T cells from MS patients resulted in increased IL-12 production compared with control T cells, consistent with the observation that CD40/CD40L

Human patients with MS have demonstrated an increase in IL-12 expression as

interactions are potent inducers of IL-12.

Elevated levels of IL-12 p70 have been detected in the synovia of RA patients compared with healthy controls (Morita et al (1998) Arthritis and Rheumatism. 41: 306-314). Cytokine messenger ribonucleic acid (mRNA) expression profile in the RA synovia identified predominantly Th1 cytokines. (Bucht et al., (1996) Clin. Exp associated with Crohn's disease (CD). Increased expression of INFy and IL-12 has been observed in the intestinal mucosa of patients with this disease (Fais et al. (1994) Linterferon Res. 14:235-238; Parronchi et al., (1997) Am. J. Path. 150:823-832; Monteleone et al., (1997) Gastroenterology. 112:1169-1178, and Berrebi et al., (1998) propria of CD patients is characteristic of a predominantly Th1 response, including greatly elevated IFNy levels (Fuss, et al., (1996) J. Immunol. 157:1261-1270).

Moreover, colon tissue sections from CD patients show an abundance of IL-12 expressing macrophages and IFNy expressing T cells (Patronchi et al. (1997) Am. J. Path. 150:823-832).

strategies have been designed to inhibit or counteract IL-12 activity. In particular, antibodies that bind to, and neutralize, IL-12 have been sought as a means to inhibit IL-12 activity. Some of the earliest antibodies were murine monoclonal antibodies (mAbs), secreted by hybridomas prepared from lymphocytes of mice immunized with IL-12 (see e.g., World Patent Application Publication No. WO 97/15327 by Strober et al.; Neurath et al. (1995) J. Exp. Med. 182:1281-1290; Duchmann et al. (1995) J. Immunol. 26:934-

Due to the role of human IL-12 in a variety of human disorders, therapeutic

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938). These murine IL-12 antibodies are limited for their use in vivo due to problems associated with administration of mouse antibodies to humans, such as short serum half life, an inability to trigger certain human effector functions and elicitation of an unwanted immune response against the mouse antibody in a human (the "human antimouse antibody" (HAMA) reaction).

In general, attempts to overcome the problems associated with use of fully-murine antibodies in humans, have involved genetically engineering the antibodies to be more "human-like." For example, chimeric antibodies, in which the variable regions of the antibody chains are murine-derived and the constant regions of the antibody chains the antibody chains are murine-derived and the constant regions of the antibody chains the antibody chains are murine-derived and the constant regions of the antibody chains are human-derived, have been prepared (Junghans, et al. (1990) Cancer Res. 50:1495-1502; Brown et al. (1991) Proc. Nail. Acad. Sci. 88:2663-2667; Kettleborough et al. (1991) Protein Engineering. 4:773-783). However, because these chimeric and humanized antibodies still retain some murine sequences, they still may elicit an unwanted immune reaction, the human anti-chimeric antibody (HACA) reaction,

especially when administered for prolonged periods.

A preferred IL-12 inhibitory agent to murine antibodies or derivatives thereof

(e.g., chimeric or humanized antibodies) would be an entirely human anti-IL-I2 antibody, since such an agent should not elicit the HAMA reaction, even if used for prolonged periods. However, such antibodies have not been described in the art and,

Summary of the Invention

therefore are still needed.

The present invention provides human antibodies that bind human IL-12. The invention also relates to the treatment or prevention of acute or chronic diseases or conditions whose pathology involves IL-12, using the human anti-IL-12 antibodies of

the invention.

In one aspect, the invention provides an isolated human antibody, or an antigen-binding portion thereof. that binds to human II. 12.

binding portion thereof, that binds to human IL-I2.

In one embodiment, the invention provides a selectively mutated human IL-I2

antibody, comprising:

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a human antibody or antigen-binding portion thereof, selectively mutated at a preferred selective mutagenesis position, contact or hypermutation position with an activity enhancing amino acid residue such that it binds to human IL-12.

In a preferred embodiment, the invention provides a selectively mutated human

L-12 antibody, comprising:
a human antibody or antigen-binding portion thereof, selectively mutated at a preferred selective mutagenesis position with an activity enhancing amino acid residue such that it binds to human IL-12.

In another preferred embodiment, the selectively mutated human IL-12 antibody

In another embodiment, the invention provides an isolated human antibody, or production of an antibody with a close to a germline immunoglobulin sequence. reactivity with other proteins or human tissues, preservation of epitope recognition, retains at least one desirable property or characteristic, e.g., preservation of non-cross another preferred embodiment, the selectively mutated human IL-12 antibody further selecting for an antibody against the same antigen using phage display technology. In affinity level is attained, the target level being improved over that attainable when antigen-binding portion thereof, is selectively mutated such that a target specificity another preferred embodiment, the selectively mutated human IL-12 antibody or preferred selective mutagenesis position, contact or hypermutation positions. In yet antibody or antigen-binding portion thereof is selectively mutated at no more than two positions. In another preferred embodiment, the selectively mutated human IL-12 more than three preferred selective mutagenesis positions, contact or hypermutation numan IL-12 antibody or antigen-binding portion thereot is selectively mutated at no enhancing amino acid residue. In another preferred embodiment, the selectively mutated selective mutagenesis position, contact or hypermutation positions with an activity or antigen-binding portion thereof is selectively mutated at more than one preferred

antigen-binding portion thereof, that binds to human IL-12 and dissociates from human IL-12 with a K_{off} rate constant of 0.1 s⁻¹ or less, as determined by surface plasmon resonance, or which inhibits phytohemagglutinin blast proliferation in an *in vitro* phytohemagglutinin blast proliferation assay (PHA assay) with an IC₅₀ of 1 x 10⁻⁶ M or less. More preferably, the isolated human antibody or an antigen-binding portion thereof, dissociates from human IL-12 with a K_{off} rate constant of 1 x 10⁻² s⁻¹ or less, or

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inhibits phytohemagglutinin blast proliferation in an *in vitro* PHA assay with an IC₅₀ of 1 x 10⁻⁷ M or less. More preferably, the isolated human antibody. or an antigen-binding portion thereof, dissociates from human IL-12 with a K_{off} rate constant of 1 x 10⁻³ s⁻¹ or less, or inhibits phytohemagglutinin blast proliferation in an *in vitro* PHA assay with an IC₅₀ of 1 x 10⁻⁸ M or less. More preferably, the isolated human antibody, or an antigenbinding portion thereof, dissociates from human IL-12 with a K_{off} rate constant of 1 x 10⁻⁹ M or less. More preferably, the isolated human antibody, or an antigenor and an intigental contains and intigental phytohemagglutinin blast proliferation in an *in vitro* PHA assay with an IC₅₀ of 1 x 10⁻⁹ M or less. More preferably, the isolated human antibody, or an antigen-binding portion thereof, dissociates from human IL-12 with a K_{off} rate constant of 1 x 10⁻⁵ s⁻¹ or less, or inhibits phytohemagglutinin blast proliferation in an *in vitro* PHA assay with an IC₅₀ of 1 x 10⁻¹⁰ M or less. Even more preferably, the isolated human antibody, or an antigen-binding portion thereof, dissociates from human IL-12 with a K_{off} rate constant of 1 x 10⁻⁵ s⁻¹ or less, or inhibits phytohemagglutinin blast proliferation in an *in vitro* PHA assay with an IC₅₀ of 1 x 10⁻¹¹ M or less.

antigen-binding portion thereof, which has the following characteristics:

a) inhibits phytohemagglutinin blast proliferation in an in vitro PHA assay

In another embodiment, the invention provides an isolated human antibody, or an

with an IC₅₀ of 1 x 10⁻⁶ M or less;

has a light chain CDR3 comprising the amino acid sequence of SEQ ID

- b) has a heavy chain CDR3 comprising the amino acid sequence of SEQ ID (O NO: 1; and
- NO: 2.

 In a preferred embodiment, the isolated human antibody, or an antigen-binding portion thereof, has a heavy chain CDR2 comprising the amino acid sequence of SEQ ID NO: 3; and has a light chain CDR1 comprising the amino acid sequence of SEQ ID NO: 4. In a preferred embodiment, the isolated human antibody, or an antigen-binding portion thereof, has a heavy chain CDR1 comprising the amino acid sequence of SEQ ID NO: 5; and has a light chain CDR1 comprising the amino acid sequence of SEQ ID NO: 6. In a preferred embodiment, the isolated human antibody, or antigen binding portion thereof, has a heavy chain variable region comprising the amino acid sequence of SEQ ID NO: 7; and has a light chain variable region comprising the amino acid sequence

sequence of SEQ ID NO: 8.

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In another embodiment, the invention provides an isolated human antibody, or an

inhibits phytohemagglutinin blast proliferation in an in vitro PHA assay antigen-binding portion thereof, which has the following characteristics:

- has a heavy chain CDR3 comprising the amino acid sequence of SEQ ID (q with an IC₅₀ of 1 x 10-9 M or less;
- NO: 9; and
- .01:0N has a light chain CDR3 comprising the amino acid sequence of SEQ ID

portion thereof, has a heavy chain CDR1 comprising the amino acid sequence of SEQ ID NO: 11; and has a light chain CDR2 comprising the amino acid sequence of SEQ ID portion thereof, has a heavy chain CDR2 comprising the amino acid sequence of SEQ In a preferred embodiment, the isolated human antibody, or an antigen-binding

- variable region comprising the amino acid sequence of SEQ ID NO: 15; and has a light NO: 14. In a preferred embodiment, the isolated human antibody has a heavy chain ID NO: 13; and has a light chain CDR1 comprising the amino acid sequence of SEQ ID NO: 12. In a preferred embodiment, the isolated human antibody, or an antigen-binding
- In another embodiment, the invention provides an isolated human antibody, or an chain variable region comprising the amino acid sequence of SEQ ID NO: 16.
- inhibits phytohemagglutinin blast proliferation in an in vitro PHA assay 50 antigen-binding portion thereof, which
- has a heavy chain CDR3 comprising the amino acid sequence of SEQ ID with an IC_{50} of 1 x 10-9 M or less;
- has a light chain CDR3 comprising the amino acid sequence of SEQ ID (c) NO: 17; and

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22. In a preferred embodiment, the isolated human antibody, or an antigen-binding ID MO: 21; and a light chain CDR1 comprising the amino acid sequence of SEQ ID MO: portion thereof, has a heavy chain CDRI comprising the amino acid sequence of SEQ 20. In a preferred embodiment, the isolated human antibody, or an antigen-binding ID MO: 19; and a light chain CDR2 comprising the amino acid sequence of SEQ ID MO: portion thereof, has a heavy chain CDR2 comprising the amino acid sequence of SEQ In a preferred embodiment, the isolated human antibody, or an antigen-binding

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lgG2, IgG3, IgG4, IgM, IgM and IgE constant regions or any allelic variation thereof as comprises a heavy chain constant region selected from the group consisting of IgOl. sequence of SEQ ID NO: 24. In a preferred embodiment, the isolated human antibody of SEQ ID NO: 23, and a light chain variable region comprising the amino acid portion thereof, has the heavy chain variable region comprising the amino acid sequence

embodiment, the isolated human antibody is a Fab fragment, or a $F(ab')_2$ fragment or a embodiment, the antibody heavy chain constant region is IgG1. In another preferred NIH Publication No. 91-3242), included herein by reference. In a more preferred Immunological Interest, Fifth Edition, U.S. Department of Health and Human Services, discussed in Kabat et al. (Kabat, E.A., et al. (1991) Sequences of Proteins of

single chain Fy fragment.

In another embodiment, the invention provides an isolated human antibody, or an

- inhibits phytohemagglutinin blast proliferation in an in vitro PHA assay antigen-binding portion thereof, which
- has a heavy chain CDR3 comprising the amino acid sequence selected with an IC_{50} of 1 x 10-9 M or less;
- has a light chain CDR3 comprising the amino acid sequence selected from the group consisting of SEQ ID NO: 404-SEQ ID NO: 469; and

from the group consisting of SEQ ID NO: 534-SEQ ID NO: 579.

preferred embodiment, the isolated human antibody comprises a heavy chain constant light chain variable region comprising the amino acid sequence of SEQ ID NO: 24. In a chain variable region comprising the amino acid sequence of SEQ ID NO: 23, and a isolated human antibody, or an antigen-binding portion thereof, comprising a the heavy consisting of SEQ ID NO: 470-SEQ ID NO: 505. In a preferred embodiment, the a light chain CDR1 comprising the amino acid sequence selected from the group sequence selected from the group consisting of SEQ ID NO: 288-SEQ ID NO: 334; and antigen-binding portion thereof, has a heavy chain CDR1 comprising the amino acid 506-SEQ ID NO: 533. In a preferred embodiment, the isolated human antibody, or an comprising the amino acid sequence selected from the group consisting of SEQ ID NO: from the group consisting of SEQ ID NO:335-SEQ ID NO: 403; and a light chain CDR2 portion thereof, has a heavy chain CDR2 comprising the amino acid sequence selected In a preferred embodiment, the isolated human antibody, or an antigen-binding 50

region, or an Fab fragment or a F(ab')2 fragment or a single chain Fy fragment as

In another embodiment, the invention provides an isolated human antibody. or an described above.

- inhibits phytohemagglutinin blast proliferation in an in vitro PHA assay antigen-binding portion thereof, which
- has a heavy chain CDR3 comprising the amino acid sequence of SEQ ID with an IC_{50} of 1 x 10-9 M or less;
- has a light chain CDR3 comprising the amino acid sequence of SEQ ID NO: 25; and
- NO: 26. (၁

acid sequence of SEQ ID NO: 32. In a preferred embodiment, the isolated human sequence of SEQ ID NO: 31, and a light chain variable region comprising the amino portion thereof, which has a heavy chain variable region comprising the amino acid 30. In a preferred embodiment, the isolated human antibody, or an antigen-binding ID NO: 29; and a light chain CDRI comprising the amino acid sequence of SEQ ID NO: portion thereof, has a heavy chain CDRI comprising the amino acid sequence of SEQ 28. In a preferred embodiment, the isolated human antibody, or an antigen-binding ID NO: 27; and a light chain CDR2 comprising the amino acid sequence of SEQ ID NO: portion thereof, has a heavy chain CDR2 comprising the amino acid sequence of SEQ. In a preferred embodiment, the isolated human antibody, or an antigen-binding

- antibody comprises a heavy chain constant region, or an Fab fragment, or a $F(ab)_2$
- In another embodiment, the invention provides an isolated human antibody, or an fragment or a single chain Fv fragment as described above.
- inhibits phytohemagglutinin blast proliferation in an in vitro PHA assay 57 antigen-binding portion thereof, which

with an IC₅₀ of 1 x 10-6 M or less;

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- or a mutant thereof having one or more amino acid substitutions at a contact position or NO: 3 and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID NO: 5, SEQ ID NO: 1, a heavy chain CDR2 comprising the amino acid sequence of SEQ ID comprises a heavy chain CDR3 comprising the amino acid sequence of
- higher than the antibody comprising a heavy chain CDR3 comprising the amino acid a hypermutation position, wherein said mutant has a koff rate no more than 10-fold

ID NO: 2; and SEQ ID NO: 3, and a heavy chain CDR1 comprising the amino acid sequence of SEQ sequence of SEQ ID NO: I, a heavy chain CDR2 comprising the amino acid sequence of

comprises a light chain CDR3 comprising the amino acid sequence of

- 4, and a light chain CDR1 comprising the amino acid sequence of SEQ ID NO: 6. SEQ ID NO: 2, a light chain CDR2 comprising the amino acid sequence of SEQ ID NO: than the antibody comprising a light chain CDR3 comprising the amino acid sequence of hypermutation position, wherein said mutant has a koff rate no more than 10-fold higher mutant thereof having one or more amino acid substitutions at a contact position or a 4, and a light chain CDR1 comprising the amino acid sequence of SEQ ID NO: 6, or a SEQ ID NO: 2, a light chain CDR2 comprising the amino acid sequence of SEQ ID NO:
- In another embodiment, the invention provides an isolated human antibody, or an
- inhibits phytohemagglutinin blast proliferation in an in vitro PHA assay antigen-binding portion thereof, which

with an IC_{50} of 1 x 10-9 M or less;

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- ID NO: 13; and SEQ ID NO: 11, and a heavy chain CDR1 comprising the amino acid sequence of SEQ sequence of SEQ ID NO: 9, a heavy chain CDR2 comprising the amino acid sequence of higher than the antibody comprising a heavy chain CDR3 comprising the amino acid or a hypermutation position, wherein said mutant has a koff rate no more than 10-fold 13, or a mutant thereof having one or more amino acid substitutions at a contact position NO: II and a heavy chain CDRI comprising the amino acid sequence of SEQ ID NO: SEQ ID NO: 9, a heavy chain CDR2 comprising the amino acid sequence of SEQ ID comprises a heavy chain CDR3 comprising the amino acid sequence of
- light chain CDR3 comprising the amino acid sequence of SEQ ID NO: 10, a light chain said mutant has a koff rate no more than 10-fold higher than the antibody comprising a selective mutagenesis position, contact position or a hypermutation position, wherein 14, or a mutant thereof having one or more amino acid substitutions at a preferred NO: 12, and a light chain CDR1 comprising the amino acid sequence of SEQ ID NO: SEQ ID NO: 10, a light chain CDR2 comprising the amino acid sequence of SEQ ID comprises a light chain CDR3 comprising the amino acid sequence of () 52

CDR2 comprising the amino acid sequence of SEQ ID NO: 12, and a light chain CDR1

comprising the amino acid sequence of SEQ ID NO: 14.

In another embodiment, the invention provides an isolated human antibody, or an

- antigen-binding portion thereof, which

 and in his provident on an in vitro PHA assay

 and in the provident of the provident
- with an IC₅₀ of 1 x 10-9 M or less;

 b) comprises a heavy chain CDR3 comprising the amino acid sequence of SEQ ID MO: 17, a heavy chain CDR1 comprising the amino acid sequence of SEQ ID MO: 19 and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID MO: 19 and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID MO: 19 and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID MO: 19 and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID MO: 19 and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID MO: 19 and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID MO: 19 and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID MO: 19 and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID MO: 19 and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID MO: 19 and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID MO: 19 and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID MO: 19 and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID MO: 19 and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID MO: 19 and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID MO: 19 and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID MO: 19 and a heavy chain CDR2 comprising the acid sequence of SEQ ID MO: 19 and a heavy chain CDR2 comprising the acid sequence of SEQ ID MO: 19 and a heavy chain CDR2 comprising the acid sequence of SEQ ID MO: 19 and a heavy chain CDR2 comprising the acid sequence of SEQ ID MO: 19 and a heavy chain CDR2 comprising the acid sequence of SEQ ID MO: 19 and a heavy chain CDR2 comprising the acid sequence of SEQ ID MO: 19 and - 21, or a mutant thereof having one or more amino acid substitutions at a preferred selective mutagenesis position, contact position or a hypermutation position, wherein said mutant has a k_{off} rate no more than 10-fold higher than the antibody comprising a heavy chain CDR3 comprising the amino acid sequence of SEQ ID NO: 19, and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID NO: 19, and a heavy chain chain CDR1 comprising the amino acid sequence of SEQ ID NO: 21; and
- comprises a light chain CDR3 comprising the amino acid sequence of SEQ ID NO: 18, a light chain CDR1 comprising the amino acid sequence of SEQ ID NO: 20, and a light chain CDR1 comprising the amino acid sequence of SEQ ID NO: 22, or a mutant thereof having one or more amino acid substitutions at preferred selective mutagenesis position, contact position or a hypermutation position, wherein said mutant has a k_{off} rate no more than 10-fold higher than the antibody comprising a light chain CDR3 comprising the amino acid sequence of SEQ ID NO: 18, a light chain

CDR2 comprising the amino acid sequence of SEQ ID NO: 20, and a light chain CDR1

antigen binding portions thereof, of the invention. A preferred isolated nucleic acid encodes the heavy chain CDR3 comprising the amino acid sequence of SEQ ID NO: 17.

The isolated nucleic acid encoding an antibody heavy chain variable region. In another embodiment, the isolated nucleic acid encodes the CDR2 of the antibody heavy chain variable region. In another variable region comprising the amino acid sequence of SEQ ID NO: 19. In another variable region comprising the amino acid sequence of SEQ ID NO: 19. In another variable region comprising the amino acid sequence of SEQ ID NO: 19. In another variable region comprising the amino acid sequence of SEQ ID NO: 19. In another variable region comprising the amino acid sequence of SEQ ID NO: 19. In another combodiment, the isolated nucleic acid encodes the CDR1 of the antibody heavy chain

comprising the amino acid sequence of SEQ ID NO: 22.

variable region comprising the amino acid sequence of SEQ ID NO: 21. In another

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embodiment, the isolated nucleic acid encodes an antibody heavy chain variable region comprising the amino acid sequence of SEQ ID NO: 23. In another embodiment, the isolated nucleic acid encodes the light chain CDR3 comprising the amino acid sequence of SEQ ID NO: 18. The isolated nucleic acid encodes the CDR2 of the region. In another embodiment, the isolated nucleic acid encodes the CDR2 of the antibody light chain variable region comprising the amino acid sequence of SEQ ID NO: 20. In another embodiment, the isolated nucleic acid encodes the CDR1 of the antibody 10.

antibody light chain variable region comprising the amino acid sequence of SEQ ID NO 20. In another embodiment, the isolated nucleic acid encodes the CDR1 of the antibody light chain variable region comprising the amino acid sequence of SEQ ID NO: 22. In another embodiment, the isolated nucleic acid encodes an antibody light chain variable region comprising the amino acid sequence of SEQ ID NO: 24.

In another embodiment, the invention provides an isolated human antibody, or an

antigen-binding portion thereof, which as proliferation in an in vitro PHA assay

with an IC₅₀ of 1 x 10-9 M or less;

b) comprises a heavy chain CDR3 comprising the amino acid sequence of SEQ ID MO: 25, a heavy chain CDR2 comprising the amino acid sequence of SEQ ID MO: 27 and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID MO: 29, or a mutant thereof having one or more amino acid substitutions at a preferred said mutant has a k_{off} rate no more than 10-fold higher than the antibody comprising a heavy chain CDR3 comprising the amino acid sequence of SEQ ID MO: 25, a heavy chain CDR2 comprising the amino acid sequence of SEQ ID MO: 27, and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID MO: 29, and a comprising the amino acid sequence of SEQ ID MO: 29, and a heavy chain comprising the amino acid sequence of SEQ ID MO: 29, and a heavy chain comprises a light chain CDR3 comprising the amino acid sequence of SEQ ID MO: 29, and

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SEQ ID NO: 26, a light chain CDR1 comprising the amino acid sequence of SEQ ID NO: 28, and a light chain CDR1 comprising the amino acid sequence of SEQ ID NO: 30, or a mutant thereof having one or more amino acid substitutions at a preferred said mutant has a k_{off} rate no more than 10-fold higher than the antibody comprising a light chain CDR3 comprising the amino acid sequence of SEQ ID NO: 26, a light chain CDR2 comprising the amino acid sequence of SEQ ID NO: 28, and a light chain CDR1 comprising the amino acid sequence of SEQ ID NO: 30.

A preferred isolated nucleic acid encodes the heavy chain CDR3 comprising the

amino acid sequence of SEQ ID NO: 25. The isolated nucleic acid encoding an antibody heavy chain variable region. In another embodiment, the isolated nucleic acid encodes the CDR2 of the antibody heavy chain variable region comprising the amino acid sequence of SEQ ID NO: 27. In another embodiment, the isolated nucleic acid encodes the CDR1 of the antibody heavy chain variable region comprising the amino acid sequence of SEQ ID NO: 29. In another embodiment, the isolated nucleic acid encodes an antibody heavy chain variable region comprising the amino acid sequence of SEQ ID NO: 26. The isolated nucleic acid encodes the light chain variable region. In another embodiment, the isolated nucleic acid encodes the CDR2 of the antibody light chain variable region the isolated nucleic acid encodes the CDR2 of the antibody light chain variable region comprising the amino acid sequence of SEQ ID NO: 28. In another embodiment, the isolated nucleic acid encodes the CDR1 of the antibody light chain variable region comprising the amino acid sequence of SEQ ID NO: 30. In another embodiment, the isolated nucleic acid encodes an antibody light chain variable region comprising the amino acid sequence of SEQ ID NO: 30. In another embodiment, the isolated nucleic acid encodes an antibody light chain variable region comprising the amino acid sequence of SEQ ID NO: 30. In another embodiment, the isolated nucleic acid encodes an antibody light chain variable region comprising the

amino acid sequence of SEQ ID NO: 32. In another aspect, the invention provides an isolated human antibody, or an

antigen-binding portion thereof, which has the following characteristics:

a) that binds to human IL-12 and dissociates from human IL-12 with a

 k_{off} rate constant of 0.1 s⁻¹or less, as determined by surface plasmon resonance, or which inhibits phytohemagglutinin blast proliferation in an *in vitro* phytohemagglutinin blast proliferation assay (PHA assay) with an IC₅₀ of 1 x 10⁻⁶M or less.

b) has a heavy chain variable region comprising an amino acid sequence

selected from a member of the V_H3 germline family, wherein the heavy chain variable region has a mutation at a preferred selective mutagenesis position, contact or hypermutation position with an activity enhancing amino acid residue.

c) has a light chain variable region comprising an amino acid sequence

selected from a member of the $V_{\lambda}I$ germline family, wherein the light chain variable region has a mutation at a preferred selective mutagenesis position, contact position or hypermutation position with an activity enhancing amino acid residue.

In another embodiment, the invention provides an isolated human antibody, or an

antigen-binding portion thereof, which has the following characteristics:

- a) that binds to human IL-12 and dissociates from human IL-12 with a
- koff rate constant of 0.1s⁻¹or less, as determined by surface plasmon resonance, or which
- inhibits phytohemagglutinin blast proliferation in an in vitro phytohemagglutinin blast proliferation assay (PHA assay) with an ${\rm IC_{50}}$ of 1 x $10^{-6}{\rm M}$ or less.
- b) has a heavy chain variable region comprising an amino acid sequence selected from the group consisting of SEQ ID NOs: 595-667, wherein the heavy chain variable region has a mutation at a preferred selective mutagenesis position, contact position or hypermutation position with an activity enhancing amino acid residue.
- c) has a light chain variable region comprising an amino acid sequence selected from the group consisting of SEQ ID NOs: 669-675, wherein the light chain variable region has a mutation at a preferred selective mutagenesis position, contact or hypermutation position with an activity enhancing amino acid residue.
- In another embodiment, the invention provides an isolated human antibody, or an antigen-binding portion thereof, which has the following characteristics:
- a) that binds to human IL-12 and dissociates from human IL-12 with a k_{off} rate constant of 0.1 s⁻¹ or less, as determined by surface plasmon resonance, or which inhibits phytohemagglutinin blast proliferation in an *in vitro* phytohemagglutinin blast proliferation assay (PHA assay) with an IC₅₀ of 1 x 10⁻⁶M or less.
- b) has a heavy chain variable region comprising the COS-3 germline amino acid sequence, wherein the heavy chain variable region has a mutation at a preferred selective mutagenesis position, contact or hypermutation position with an activity enhancing amino acid residue.

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acid sequence, wherein the light chain variable region has a mutation at a preferred selective mutagenesis position, contact or hypermutation position with an activity enhancing amino acid residue.

c) has a light chain variable region comprising the DPL8 germline amino

In another embodiment, the invention provides an isolated human antibody, or an antigen-binding portion thereof, which has the following characteristics:

a) that binds to human IL-12 and dissociates from human IL-12 with a $k_{\rm off}$ rate constant of 0.1 s⁻¹ or less, as determined by surface plasmon resonance, or which inhibits phytohemagglutinin blast proliferation in an *in vitro* phytohemagglutinin blast proliferation as an IC₅₀ of 1 x 10⁻⁶M or less.

selected from a member of the $V_{\rm H}3$ germline family, wherein the heavy chain variable region comprises a CDR2 that is structurally similar to CDR2s from other $V_{\rm H}3$ germline family members, and a CDR1 that is structurally similar to CDR1s from other $V_{\rm H}3$ germline family members, and wherein the heavy chain variable region has a mutation at a preferred selective mutagenesis position, contact or hypermutation position with an

activity enhancing amino acid residue;
c) has a light chain variable region comprising an amino acid sequence

selected from a member of the $V_{\lambda}I$ germline family, wherein the light chain variable region comprises a CDR2 that is atructurally similar to CDR2s from other $V_{\lambda}I$ germline family members, and a CDR1 that is atructurally similar to CDR1s from other $V_{\lambda}I$ germline family members, and wherein the light chain variable region has a mutation at a preferred selective mutagenesis position, contact or hypermutation position with an activity enhancing amino acid residue.

activity enhancing amino acid residue.

In a preferred embodiment, the isolated human antibody, or antigen binding

portion thereof, has a mutation in the heavy chain CDR3. In another preferred embodiment, the isolated human antibody, or antigen binding portion thereof, has a mutation in the light chain CDR3. In another embodiment, the isolated human antibody, or antigen binding portion thereof, has a mutation in the light chain CDR2. In another preferred embodiment, the isolated human antibody, or antigen binding portion thereof, has a mutation in the light chain CDR2. In another preferred embodiment, the isolated human antibody, or antigen binding portion thereof, has a mutation in the light chain the isolated binding portion thereof, has a mutation in the heavy chain cDR1. In another preferred embodiment, the isolated human antibody, or antigen binding portion thereof, has a mutation in the light chain CDR1.

In another aspect, the invention provides recombinant expression vectors

so carrying the antibody-encoding nucleic acids of the invention, and host cells into which

such vectors have been introduced, are also encompassed by the invention, as are

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methods of making the antibodies of the invention by culturing the host cells of the invention.

In another aspect, the invention provides an isolated human antibody, or antigenbinding portion thereof, that neutralizes the activity of human IL-12, and at least one additional primate IL-12 selected from the group consisting of baboon IL-12, marmoset IL-12, chimpanzee IL-12, cynomolgus IL-12 and rhesus IL-12, but which does not

neutralize the activity of the mouse IL-12.

In another aspect, the invention provides a pharmaceutical composition comprising the antibody or an antigen binding portion thereof, of the invention and a

In another aspect, the invention provides a composition comprising the antibody or an antigen binding portion thereof, and an additional agent, for example, a therapeutic agent.

In another aspect, the invention provides a method for inhibiting human IL-12 activity comprising contacting human IL-12 with the antibody of the invention, e.g.,

1695, such that human IL-12 activity is inhibited.

In another aspect, the invention provides a method for inhibiting human IL-12 activity is a human subject suffering from a disorder in which IL-12 activity is

activity in a human subject suffering from a disorder in which IL-12 activity is detrimental, comprising administering to the human subject the antibody of the invention, e.g., 1695, such that human IL-12 activity in the human subject is inhibited.

The disorder can be, for example, Crohn's disease, multiple seletosis or rheumatoid

arthritis.

In another aspect, the invention features a method for improving the activity of an antibody, or an antigen binding portion thereof, to attain a predetermined target

25 activity, comprising:

antibodies, or antigen binding portions thereof;

pharmaceutically acceptable carrier.

- a) providing a parent antibody a antigen-binding portion thereof;

 consisting of H30, H31, H31B, H32, H33, H52, H56, H58, L30, L31, L32, L50, L91,

 consisting of H30, H31, H31B, H32, H33, H52, H56, H58, L30, L31, L32, L50, L91,
- 30 c) individually mutating the selected preferred selective mutagenesis position to at least two other amino acid residues to hereby create a first panel of mutated

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antigen binding portions thereof to determined if mutation of a single selective d) evaluating the activity of the first panel of mutated antibodies, or

e) combining in a stepwise fashion, in the parent antibody, or antigen binding predetermined target activity or a partial target activity; mutagenesis position produces an antibody or antigen binding portion thereof with the

combination antibodies, or antigen binding portions thereof. portion thereof, individual mutations shown to have an improved activity, to form

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g) if steps d) or f) do not result in an antibody or antigen binding portion thereof thereof have the predetermined target activity or a partial target activity. portions thereof to determined if the combination antibodies, or antigen binding portions f) evaluating the activity of the combination antibodies, or antigen binding

H53, H54, H95, H96, H97, H98, L30A and L96 are mutated to at least two other amino activity, additional amino acid residues selected from the group consisting of H35, H50, having the predetermined target activity, or result an antibody with only a partial

acid residues to thereby create a second panel of mutated antibodies or antigen-binding

h) evaluating the activity of the second panel of mutated antibodies or antigen portions thereof;

and L96 results an antibody or antigen binding portion thereof, having the predetermined selected from the group consisting of H35, H50, H54, H95, H95, H97, H98, L30A binding portions thereof, to determined if mutation of a single amino acid residue

i) combining in stepwise fashion in the parent antibody, or antigen-binding target activity or a partial activity; .

form combination antibodies, or antigen binding portions thereof; portion thereof, individual mutations of step g) shown to have an improved activity, to

j) evaluating the activity of the combination antibodies or antigen binding 57

k) if steps h) or j) do not result in an antibody or antigen binding portion thereof portions thereof have the predetermined target activity or a partial target activity; portions thereof, to determined if the combination antibodies, or antigen binding

third panel of mutated antibodies or antigen binding portions thereof; H52B and L31A are mutated to at least two other amino acid residues to thereby create a activity, additional amino acid residues selected from the group consisting of H33B, having the predetermined target activity, or result in an antibody with only a partial

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l) evaluating the activity of the third panel of mutated antibodies or antigen binding portions thereof, to determine if a mutation of a single amino acid residue selected from the group consisting of H33B, H52B and L31A resulted in an antibody or antigen binding portion thereof, having the predetermined target activity or a partial activity;

- m) combining in a stepwise fashion in the parent antibody. or antigen binding portion thereof, individual mutation of step k) shown to have an improved activity, to form combination antibodies, or antigen binding portions, thereof;
- n) evaluating the activity of the combination antibodies or antigen-binding portions thereof, to determine if the combination antibodies, or antigen binding portions thereof have the predetermined target activity to thereby produce an antibody or antigen binding portions.

binding portion thereof with a predetermined target activity.

In another aspect, the invention provides a method for improving the activity of

an antibody, or antigen-binding portion thereof, comprising:

- a) providing a parent antibody or antigen-binding portion thereof;

 b) selecting a preferred selective mutagenesis position, contact or hypermutation position within a complementarity determining region (CDR) for mutation, thereby
- position within a complementarity determining region (CDR) for mutation, identifying a selected preferred selective mutagenesis position, contact or hypermutation position;
- c) individually mutating said selected preferred selective mutagenesis position,
 contact or hypermutation position to at least two other amino acid residues to thereby
 create a panel of mutated antibodies, or antigen-binding portions thereof;
 d) evaluating the activity of the panel of mutated antibodies, or antigen-binding

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- portions thereof, relative to the parent antibody or antigen-binding portion thereof;

 e) repeating steps b) through d) for at least one other contact or hypermutation position;
- f) combining, in the parent antibody, or antigen-binding portion thereof, individual mutations shown to have improved activity, to form combination antibodies, or antigen-binding portions thereof; and

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g) evaluating the activity of the combination antibodies, or antigen-binding portion thereof; until an antibody, or antigen-binding portion thereof, with an improved activity, relative to the parent antibody, or antigen-binding portion thereof, is obtained.

In one embodiment, the invention provides a method for improving the activity of an antibody. or antigen-binding portion thereof, comprising:

a) providing a recombinant parent antibody or antigen-binding portion thereof.

a) providing a recombinant parent antibody or antigen-binding portion thereof; that was obtained by selection in a phage-display system but whose activity is not

further improved by mutagenesis in said phage-display system;

b) selecting a preferred selective mutagenesis position, contact or hypermutation

position within a complementarity determining region (CDR) for mutation, thereby identifying a selected contact or hypermutation position;

c) individually mutating said selected preferred selective mutagenesis position,

contact or hypermutation position to at least two other amino acid residues to thereby create a panel of mutated antibodies, or antigen-binding portions thereof, and expressing

said panel in a non-phage display system;

d) evaluating the activity of the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof;

e) repeating steps b) through d) for at least one other contact or hypermutation

20 position;
f) combining, in the parent antibody, or antigen-binding portion thereof,

individual mutations shown to have improved activity, to form combination antibodies, or antigen-binding portions thereof; and
g) evaluating the activity of the combination antibodies, or antigen-binding

portions thereof, relative to the parent antibody or antigen-binding portion thereof; until an antibody, or antigen-binding portion thereof, with an improved activity, relative to the parent antibody, or antigen-binding portion thereof, is obtained.

In a preferred embodiment, the contact positions are selected from the group

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In a more preferred embodiment, the contact positions are selected from the group H30, H31, H31B, H32, H33, H52, H56, H58, L30, L31, L32, L50, L91, L92, L93, L94. selected from the preferred selective mutagenesis positions from the group consisting of L53 and L93. In a more preferred embodiment the residues for selective mutagenesis are

In another embodiment, the invention provides a method for improving the consisting of L50 and L94.

activity of an antibody, or antigen-binding portion thereof, comprising:

- a) providing a recombinant parent antibody or antigen-binding portion thereof;
- b) selecting a preferred selective mutagenesis position, contact or hypermutation
- position within a complementarity determining region (CDR) for mutation, thereby
- c) individually mutating said selected preferred selective mutagenesis position, identifying a selected contact or hypermutation position;
- create a panel of mutated antibodies, or antigen-binding portions thereof and expressing contact or hypermutation position to at least two other amino acid residues to thereby
- d) evaluating the activity of the panel of mutated antibodies, or antigen-binding said panel in an appropriate expression system;
- thereby identifying an activity enhancing amino acid residue; portions thereof, relative to the parent antibody or antigen-binding portion thereof
- other property or characteristics, wherein the property or characteristic is one that needs thereof, relative to the parent antibody or antigen-binding portion thereof for at least one e) evaluating the panel of mutated antibodies, or antigen-binding portions
- binding portion thereof, is obtained. least one retained property or characteristic, relative to the parent antibody, or antigenuntil an antibody, or antigen-binding portion thereof, with an improved activity and at to be retained in the antibody;

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recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 crossreactivity with other proteins or human tissues, 2) preservation of epitope L94 and L96 and the other characteristic is selected from 1) preservation of non-H62' H66' H61' H68' H101' F30' F31' F35' F34' F20' F25' F23' F22' F61' F65' F63' consisting of H30, H31, H31B, H32, H33, H35, H50, H52, H52A, H53, H54, H56, H58, In a preferred embodiment, the contact positions are selected from the group

heterodimer preventing binding interference from free, soluble p40 and/or 3) to produce

further improved by mutagenesis in said phage-display system; that was obtained by selection in a phage-display system but whose activity cannot be a) providing a recombinant parent antibody or antigen-binding portion thereof; activity of an antibody, or antigen-binding portion thereof, comprising: In another embodiment of the invention provides a method for improving the with a close to germline immunoglobulin sequence. preventing binding interference from free, soluble p40 and/or 3) to produce an antibody recognizing p40 epitope preferably in the context of the p70 p40/p35 hererodimer with other proteins or human tissues, 2) preservation of epitope recognition, i.e. L94 and the other characteristic is selected from 1) preservation of non-crossreactivity embodiment, the contact positions are selected from the group consisting of L50 and an antibody with a close to germline immunoglobulin sequence. In a more preferred heterodimer preventing binding interference from free, soluble p40 and/or 3) to produce recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 crossreactivity with other proteins or human tissues, 2) preservation of epitope L91, L92, L93, L94 and the other characteristic is selected from 1) preservation of nongroup consisting of H30, H31, H31B, H32, H32, H56, H56, H58, L30, L31, L32, L50, mutagenesis are selected from the preferred selective mutagenesis positions from the immunoglobulin sequence. In a more preferred embodiment the residues for selective from free, soluble p40 and/or 3) to produce an antibody with a close to germline preferably in the context of the p70 p40/p35 heterodimer preventing binding interference or human tissues, 2) preservation of epitope recognition. i.e. recognizing p40 epitope characteristic is selected from 1) preservation of non-crossreactivity with other proteins H31' H31B' H35' H25' H26' H28' L30, L31, L32, L53 and L93 and the other embodiment, the hypermutation positions are selected from the group consisting of H30. an antibody with a close to germline immunoglobulin sequence. In another preferred

identifying a selected preferred selective mutagenesis position, contact or hypermutation

b) selecting a preferred selective mutagenesis position, contact or hypermutation

position within a complementarity determining region (CDR) for mutation, thereby

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said panel in a non-phage display system; create a panel of mutated antibodies, or antigen-binding portions thereof, and expressing contact or hypermutation position to at least two other amino acid residues to thereby c) individually mutating said selected preferred selective mutagenesis position,

portions thereof, relative to the parent antibody or antigen-binding portion thereof d) evaluating the activity of the panel of mutated antibodies. or antigen-binding

e) evaluating the panel of mutated antibodies, or antigen-binding portions thereby identifying an activity enhancing amino acid residue;

antibody, or antigen-binding portion thereof, is obtained. activity and at least one retained property or characteristic, relative to the parent to be retained, until an antibody, or antigen-binding portion thereof, with an improved other property or characteristic, wherein the property or characteristic is one that needs thereof, relative to the parent antibody or antigen-binding portion thereof for at least one

f) repeating steps a) through e) for at least one other preferred selective

mutagenesis position, contact or hypermutation position;

antigen-binding portions thereof; and and at least on retained property or characteristic, to form combination antibodies, or two individual activity enhancing amino acid residues shown to have improved activity g) combining, in the parent antibody, or antigen-binding portion thereof, at least

h) evaluating the activity of the combination antibodies, or antigen-binding

binding portion thereof, is obtained. least one retained property or characteristic, relative to the parent antibody, or antigenuntil an antibody, or antigen-binding portion thereof, with an improved activity and at portions thereof, relative to the parent antibody or antigen-binding portion thereof;

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an antibody with a close to germline immunoglobulin sequence. In another preferred heterodimer preventing binding interference from free, soluble p40 and/or 3) to produce recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 crossreactivity with other proteins or human tissues, 2) preservation of epitope L94 and L96 and the other characteristic is selected from 1) preservation of non-H92' H96' H61' H68' H101' F30' F31' F35' F34' F20' F25' F23' F22' F61' F65' F63' consisting of H30, H31, H31B, H32, H33, H36, H50, H52, H52A, H53, H54, H56, H58, In a preferred embodiment, the contact positions are selected from the group 52

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preferably in the context of the p70 p40/p35 heterodimer preventing binding interference or human tissues, 2) preservation of epitope recognition, i.e. recognizing p40 epitope characteristic is selected from 1) preservation of non-crossreactivity with other proteins H31, H31B, H32, H52, H56, H58, L30, L31, L32, L53 and L93 and the other embodiment, the hypermutation positions are selected from the group consisting of H30,

- L91, L92, L93, L94 and the other characteristic is selected from 1) preservation of nongroup consisting of H30, H31, H31B, H32, H33, H52, H56, H58, L30, L31, L32, L50, mutagenesis are selected from the preferred selective mutagenesis positions from the immunoglobulin sequence. In a more preferred embodiment the residues for selective from free, soluble p40 and/or 3) to produce an antibody with a close to germline
- recognizing p40 epitope preferably in the context of the p70 p40/p35 heterodimer with other proteins or human tissues, 2) preservation of epitope recognition, i.e. L94 and the other characteristic is selected from 1) preservation of non-crossreactivity embodiment, the contact positions are selected from the group consisting of L50 and an antibody with a close to germline immunoglobulin sequence. In a more preferred heterodimer preventing binding interference from free, soluble p40 and/or 3) to produce recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 crossreactivity with other proteins or human tissues, 2) preservation of epitope
- with a close to germline immunoglobulin sequence. preventing binding interference from free, soluble p40 and/or 3) to produce an antibody
- activity of an antibody, or antigen-binding portion thereof, comprising: In another embodiment, the invention provides a method for improving the
- further improved by mutagenesis in said phage-display system; that was obtained by selection in a phage-display system but whose activity cannot be a) providing a recombinant parent antibody or antigen-binding portion thereof;
- hypermutation position; determining region (CDR) for mutation, thereby identifying a selected contact or b) selecting a contact or hypermutation position within a complementarity
- antigen-binding portions thereof, and expressing said panel in a non-phage display least two other amino acid residues to thereby create a panel of mutated antibodies, or c) individually mutating said selected contact or hypermutation position to at

system;

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d) evaluating the activity of the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereby identifying an activity enhancing amino acid residue;

e) evaluating the panel of mutated antibodies, or antigen-binding portions

thereof, relative to the parent antibody or antigen-binding portion thereof for at least one other property or characteristics, wherein the property or characteristic is one that needs to be retained;

to be retained; until an antibody, or antigen-binding portion thereof, with an improved activity and at least one retained property or characteristic, relative to the parent antibody, or antigen-binding portion thereof, is obtained.

In a preferred embodiment, the contact positions are selected from the group consisting of H30, H31, H31B, H32, H33, H35, H50, H52, L53, L53, L53, L54, H56, H58, H95, H97, H98, H101, L30, L31, L32, L34, L50, L52, L53, L53, L53, L91, L92, L93, L94 and L96 and the other characteristic is selected from 1) preservation of non-recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 heterodimer preventing binding interference from free, soluble p40 and/or 3) to produce an antibody with a close to germline immunoglobulin sequence. In another preferred embodiment, the hypermutation positions are selected from the group consisting of H30, embodiment, the hypermutation positions are selected from the group consisting of H30, embodiment, the hypermutation positions are selected from the group consisting of H30, embodiment, the hypermutation positions are selected from the group consisting of H30, H31, H31B, H32, H52, H56, H58, L30, L31, L32, L53 and L93 and the other

characteristic is selected from 1) preservation of non-crossreactivity with other proteins or human tissues, 2) preservation of epitope recognition, i.e. recognizing p40 epitope from free, soluble p40 and/or 3) to produce an antibody with a close to germline from free, soluble p40 and/or 3) to produce an antibody with a close to germline mutagenesis are selected from the preferred embodiment the residues for selective mutagenesis are selected from the preferred selective mutagenesis positions from the group consisting of H30, H31, H31B, H32, H33, H52, H56, H58, L30, L31, L32, L50, L91, L92, L93, L94 and the other characteristic is selected from 1) preservation of noncrossreactivity with other proteins or human tissues, 2) preservation of epitope crossreactivity with other proteins or human tissues, 2) preservation of p40/p35 recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 and other proteins of the p70 p40/p35

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preventing binding interference from free, soluble p40 and/or 3) to produce an antibody recognizing p40 epitope preferably in the context of the p70 p40/p35 heterodimer with other proteins or human tissues, 2) preservation of epitope recognition, i.e. L94 and the other characteristic is selected from 1) preservation of non-crossreactivity embodiment, the contact positions are selected from the group consisting of L50 and

- with a close to germline immunoglobulin sequence.
- In another embodiment, the invention provides a method for improving the
- that was obtained by selection in a phage-display system but whose activity cannot be a) providing a recombinant parent antibody or antigen-binding portion thereof; activity of an antibody, or antigen-binding portion thereof, comprising:
- b) selecting a preferred selective mutagenesis position, contact or hypermutation further improved by mutagenesis in said phage-display system;
- ;noitizoq identifying a selected preferred selective mutagenesis position contact or hypermutation position within a complementarity determining region (CDR) for mutation, thereby
- said panel in a non-phage display system; create a panel of mutated antibodies, or antigen-binding portions thereof, and expressing contact or hypermutation position to at least two other amino acid residues to thereby individually mutating said selected preferred selective mutagenesis position,
- thereby identifying an activity enhancing amino acid residue; portions thereof, relative to the parent antibody or antigen-binding portion thereof d) evaluating the activity of the panel of mutated antibodies, or antigen-binding

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e) evaluating the panel of mutated antibodies, or antigen-binding portions

- f) repeating steps a) through e) for at least one other preferred selective antibody, or antigen-binding portion thereof, is obtained. activity and at least one retained property or characteristic, relative to the parent to be retained, until an antibody, or antigen-binding portion thereof, with an improved other property or characteristic, wherein the property or characteristic is one that needs thereof, relative to the parent antibody or antigen-binding portion thereof for at least one
- mutagenesis position, contact or hypermutation position;

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g) combining, in the parent antibody, or antigen-binding portion thereof, at least two individual activity enhancing amino acid residues shown to have improved activity and at least on retained other characteristic, to form combination antibodies, or antigen-binding portions thereof; and

h) evaluating the activity of the combination antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof, with an improved activity and at least one retained property or characteristic, relative to the parent antibody, or antigenleast one retained property or characteristic, relative to the parent antibody, or antigenleast one retained property or characteristic, relative to the parent antibody, or antigenleast one retained property or characteristic, relative to the parent antibody, or antigenleast one retained property or characteristic, relative to the parent antibody, or antigenleast one retained property or characteristic.

In a preferred embodiment, the contact positions are selected from the group

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embodiment, the contact positions are selected from the group consisting of L50 and an antibody with a close to germline immunoglobulin sequence. In a more preferred heterodimer preventing binding interference from free, soluble p40 and/or 3) to produce recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 crossreactivity with other proteins or human tissues, 2) preservation of epitope L91, L92, L93, L94 and the other characteristic is selected from 1) preservation of nongroup consisting of H30, H31, H31B, H32, H33, H52, H56, H58, L30, L31, L32, L50, mutagenesis are selected from the preferred selective mutagenesis positions from the immunoglobulin sequence. In a more preferred embodiment the residues for selective from free, soluble p40 and/or 3) to produce an antibody with a close to germline preferably in the context of the p70 p40/p35 heterodimer preventing binding interference or human tissues, 2) preservation of epitope recognition, i.e. recognizing p40 epitope characteristic is selected from 1) preservation of non-crossreactivity with other proteins H31, H31B, H32, H52, H56, H58, L30, L31, L32, L53 and L93 and the other embodiment, the hypermutation positions are selected from the group consisting of H30, an antibody with a close to germline immunoglobulin sequence. In another preferred heterodimer preventing binding interference from free, soluble p40 and/or 3) to produce recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 crossreactivity with other proteins or human tissues, 2) preservation of epitope L94 and L96 and the other characteristic is selected from 1) preservation of non-H92' H96' H91' H98' H101' C30' C31' C37' C34' C20' C27' C23' C22' C31' C63' consisting of H30, H31, H31B, H32, H33, H35, H50, H52, H52A, H53, H54, H56, H58,

with a close to germline immunoglobulin sequence. preventing binding interference from free, soluble p40 and/or 3) to produce an antibody recognizing p40 epitope preferably in the context of the p70 p40/p35 heterodimer with other proteins or human tissues. 2) preservation of epitope recognition, i.e. L94 and the other characteristic is selected from 1) preservation of non-crossreactivity

- activity of an antibody, or antigen-binding portion thereof, comprising: In another embodiment, the invention provides a method for improving the
- a) providing a parent antibody or antigen-binding portion thereof;
- H23' H24' H26' H28' H62' H62' H64' H64' H101' F30' F31' F35' F20' F25' F23' (CDR) for mutation other than H30, H31, H31B, H32, H33, H35, H50, H52, H52A, b) selecting an amino acid residue within a complementarity determining region
- residues to thereby create a panel of mutated antibodies, or antigen-binding portions c) individually mutating said selected position to at least two other amino acid F22' F31' F35' F32' F34 and E36:
- e) evaluating the panel of mutated antibodies, or antigen-binding portions thereby identifying an activity enhancing amino acid residue; portions thereof, relative to the parent antibody or antigen-binding portion thereof d) evaluating the activity of the panel of mutated antibodies, or antigen-binding
- thereof, relative to the parent antibody or antigen-binding portion thereof, for changes in
- Preferably, the other characteristic or property is selected from 1) preservation of to the parent antibody, or antigen-binding portion thereof, is obtained. until an antibody, or antigen-binding portion thereof, with an improved activity, relative at least one other property or characteristic;
- an antibody with a close to germline immunoglobulin sequence heterodimer preventing binding interference from free, soluble p40 and/or 3) to produce recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 non-crossreactivity with other proteins or human tissues, 2) preservation of epitope
- activity of an antibody, or antigen-binding portion thereof, comprising: 30 In another embodiment, the invention provides a method for improving the

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thereof;

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a) providing a parent antibody or antigen-binding portion thereof;

H23' H24' H26' H28' H62' H66' H67' H68' H101' L30, L31, L32, L34, L50, L52, L53, (CDR) for mutation other than H30, H31, H31B. H32, H33, H35. H50, H52, H52A, b) selecting an amino acid residue within a complementarity determining region

c) individually mutating said selected position to at least two other amino acid ς F22' F61' F65' F63' F64 and F66:

thereof; residues to thereby create a panel of mutated antibodies, or antigen-binding portions

portions thereof, relative to the parent antibody or antigen-binding portion thereof, d) evaluating the activity of the panel of mutated antibodies, or antigen-binding

neither the position selected under b) nor a position at H30, H31, H31B, H32, H33, e) repeating steps b) through d) for at least one other CDR position which is

thereby identifying an activity enhancing amino acid residue;

T35' T34' T20' T25' T23' T22' T31' T35' T33' T34 and L36; H32' H20' H25' H254' H23' H24' H26' H28' H62' H66' H61' H68' H101' F30' F31'

g) evaluating the activity of the combination antibodies, or antigen-binding to form combination antibodies, or antigen-binding portions thereof; and two individual activity enhancing amino acid residues shown to have improved activity, f) combining, in the parent antibody, or antigen-binding portion thereof, at least

thereof, with an improved activity, relative to the parent antibody, or antigen-binding antibody or antigen-binding portion thereof until an antibody, or antigen-binding portion portions thereof with two activity enhancing amino acid residues, relative to the parent

In another embodiment, the invention provides a method for improving the portion thereof, is obtained.

a) providing a recombinant parent antibody or antigen-binding portion thereof; 52 activity of an antibody, or antigen-binding portion thereof, comprising:

further improved by mutagenesis in said phage-display system; that was obtained by selection in a phage-display system but whose activity cannot be

H23' H24' H26' H28' H62' H62' H64' H68' H101' F30' F31' F35' F34' F20' F25' F23' 30 (CDR) for mutation other than H30, H31, H31B, H32, H33, H35, H50, H52, H52A, b) selecting an amino acid residue within a complementarity determining region

L55, L91, L92, L93, L94 and;

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c) individually mutating said selected contact or hypermutation position to at least two other amino acid residues to thereby create a panel of mutated antibodies, or antigen-binding portions thereof, and expressing said panel in a non-phage display

system:

d) evaluating the activity of the panel of mutated antibodies. or antigen-binding

portions thereof, relative to the parent antibody or antigen-binding portion thereof thereby identifying an activity enhancing amino acid residue;

e) evaluating the panel of mutated antibodies, or antigen-binding portions

thereof, relative to the parent antibody or antigen-binding portion thereof, for changes in at least one other property or characteristic until an antibody, or antigen-binding portion thereof, with an improved activity, relative to the parent antibody. or antigen-binding

portion thereof, is obtained.

Preferably, the other characteristic or property is selected from 1) preservation of

Preferably, the other characteristic or property is selected from 1) preservation of non-crossreactivity with other proteins or human tissues, 2) preservation of epitope

non-crossreactivity with other proteins or human tissues, 2) preservation of epitope recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 heterodimer preventing binding interference from free, soluble p40 and/or 3) to produce

An antibody with a close to germline immunoglobulin sequence.

In another embodiment, the invention provides a method for improving the

activity of an antibody, or antigen-binding portion thereof, comprising:

a) providing a parent antibody or antigen-binding portion thereof that was
obtained by selection in a phage-display system but whose activity cannot be further

improved by mutagenesis in said phage-display system;

b) selecting an amino acid residue within a complementarity determining region

- (CDR) for mutation other than H30, H31, H31B, H32, H33, H35, H50, H52, H52A, H53, H54, H56, H58, H95, H97, H98, H101, L30, L31, L32, L34, L50, L52, L53,
- L55, L91, L92, L93, L94 and L96;
 c) individually mutating said selected position to at least two other amino acid
- c) individually mutating said selected position to at least two other amino acid tresidues to thereby create a panel of mutated antibodies, or antigen-binding portions thereof and expression in a non-phage display system;
- mereot and expression in a non-phage display system,

 d) evaluating the activity of the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof

thereby identifying an activity enhancing amino acid residue;

e) repeating steps b) through d) for at least one other position within the CDR which is neither the position selected under b) nor a position at H30, H31, H31B, H32, H33, H35, H50, H52, H52A, H53, H54, H56, H58, H95, H96, H97, H98, H101, L30, L31, L32, L34, L50, L52, L53, L55, L91, L92, L93, L94;

f) combining, in the parent antibody, or antigen-binding portion thereof, at least two individual activity enhancing amino acid residues shown to have improved activity, to form combination antibodies, or antigen-binding portions thereof; and

g) evaluating the activity and other property or characteristic of the combination antibodies, or antigen-binding portions thereof with two activity enhancing amino acid residues, relative to the parent antibody or antigen-binding portion thereof, with an improved activity, relative to the parent antibody, or antigen-binding portion thereof, is obtained.

Preferably, the other characteristic or property is selected from 1) preservation of

non-crossreactivity with other proteins or human tissues, 2) preservation of epitope recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 heterodimer preventing binding interference from free, soluble p40 and/or 3) to produce an antibody with a close to germline immunoglobulin sequence.

In another embodiment, the invention provides a method for improving the

activity of an antibody, or antigen-binding portion thereof, comprising:

a) providing a parent antibody or antigen-binding portion thereof;

b) selecting an amino acid residue within a complementarity determining region

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(CDR) for mutation other than H30, H31, H31B, H32, H33, H35, H36, L50, L52, L53, L53, H54, H56, H58, H95, H97, H98, H101, L30, L31, L32, L34, L50, L52, L53,

c) individually mutating said selected position to at least two other amino acid residues to thereby create a panel of mutated antibodies, or antigen-binding portions thereof;

d) evaluating the activity of the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof,
 thereby identifying an activity enhancing amino acid residue;

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relative to the parent antibody or antigen-portion thereof, for changes in at least one e) evaluating the panel of mutated antibodies or antigen-binding portions thereof,

H32' H20' H27' H27∀' H23' H24' H26' H28' H62' H62' H64' H68' H101' 厂31' neither the position selected under b) nor a position at H30, H31, H31B, H32, H33, f) repeating steps b) through e) for at least one other CDR position which is other property or characteristic;

g) combining, in the parent antibody, or antigen-binding portion thereof, at least L32, L34, L50, L52, L53, L55, L91, L92, L93, L94 and L96;

antibodies, or antigen-binding portions thereof; and and not affecting at least one other property or characteristic, to form combination two individual activity enhancing amino acid residues shown to have improved activity

binding portion thereof until an antibody, or antigen-binding portion thereof, with an activity enhancing amino acid residues, relative to the parent antibody or antigenproperty of the combination antibodies, or antigen-binding portions thereof with two h) evaluating the activity and the retention of at least one other characteristic or

parent antibody, or antigen-binding portion thereof, is obtained. improved activity and at least one retained property or characteristic, relative to the

In another embodiment the invention provides a method to improve the affinity

obtained by selection in a phage-display system but whose activity cannot be further a) providing a parent antibody or antigen-binding portion thereof that was 50 of an antibody or antigen-binding portion thereof, comprising:

b) selecting an amino acid residue within a complementarity determining region improved by mutagenesis in said phage-display system;

H23' H24' H26' H28' H32' H30' H3L' H38' H101' F30' F31' F35' F34' F20' F25' F23' 52 (CDR) for mutation other than H30, H31, H31B, H32, H33, H35, H50, H52, H52A,

thereof and expression in a non-phage display system; residues to thereby create a panel of mutated antibodies, or antigen-binding portions c) individually mutating said selected position to at least two other amino acid F22' F31' F35' F33' F34 and E36;

thereby identifying an activity enhancing amino acid residue; portions thereof, relative to the parent antibody or antigen-binding portion thereof d) evaluating the activity of the panel of mutated antibodies, or antigen-binding

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- e) evaluating the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion at least one other characteristic or property until an antibody, or antigen-binding portion thereof, with an improved activity, relative to the parent antibody, or antigen-binding
- portion thereof, is obtained.

 In another embodiment, the invention provides a method for improving the
- activity of an antibody, or antigen-binding portion thereof, comprising:

 a) providing a parent antibody or antigen-binding portion thereof;
- b) selecting an amino acid residue within a complementarity determining region
- b) selecting an amino acid residue within a complementarity determining region (CDR) for mutation at a position other than H30, H31, H31B, H32, H33, H35, H50,
- H52, H52A, H53, H54, H56, H58, H95, H96, H97, H98, H101, L30, L31, L32, L34,
- L50, L52, L53, L53, L51, L92, L93. L94 and L96;
 c) individually mutating said selected position to at least two other amino acid residues to thereby create a panel of mutated antibodies, or antigen-binding portions
- thereof;
 d) evaluating the activity of the panel of mutated antibodies, or antigen-binding
- portions thereof, relative to the parent antibody or antigen-binding portion thereof, thereby identifying an activity enhancing amino acid residue;
- e) evaluating the panel of mutated antibodies or antigen-binding portions thereof, relative to the parent antibody or antigen-portion thereof, for changes in at least one
- other property or characteristic;

 f) repeating steps b) through e) for at least one other CDR position which is neither the position selected under b) nor a position at H30, H31, H31B, H32, H33, H35, H50, H52, H52A, H53, H54, H56. H58, H95, H96, H97, H98, H101, L30, L31, L32,
- g) combining, in the parent antibody, or antigen-binding portion thereof, at least two individual activity enhancing amino acid residues shown to have improved activity but not affecting at least one other property or characteristic, to form combination antibodies, or antigen-binding portions thereof with at least one retained property or

L34, L50, L52, L53, L55, L91, L92, L93, L94 and L96;

characteristic; and

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improved activity and at least one retained property or characteristic. relative to the binding portion thereof until an antibody, or antigen-binding portion thereof, with an two activity enhancing amino acid residues, relative to the parent antibody or antigencharacteristic of the combination antibodies, or antigen-binding portions thereof with h) evaluating the activity and the retention of at least one property of

parent antibody, or antigen-binding portion thereof, is obtained.

In another embodiment, the invention provides a method for improving the an antibody with a close to germline immunoglobulin sequence heterodimer preventing binding interference from free, soluble p40 and/or 3) to produce recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 non-crossreactivity with other proteins or human tissues, 2) preservation of epitope Preferably, the other characteristic or property is selected from 1) preservation of

activity of an antibody, or antigen-binding portion thereof, without affecting other

- a) providing a parent antibody or antigen-binding portion thereof; characteristics, comprising:
- L55, L91, L92, L93, L94 and L96; H23' H24' H26' H28' H62' H62' H64' H64' H101' F30' F31' F37' F20' F27' F23' (CDR) for mutation other than H30, H31, H31B, H32, H33, H35, H50, H52, H52A, b) selecting an amino acid residue within a complementarity determining region
- thereof; residues to thereby create a panel of mutated antibodies, or antigen-binding portions c) individually mutating said selected position to at least two other amino acid
- thereby identifying an activity enhancing amino acid residue; portions thereof, relative to the parent antibody or antigen-binding portion thereof d) evaluating the activity of the panel of mutated antibodies, or antigen-binding
- to the parent antibody, or antigen-binding portion thereof, is obtained. thereof, with an improved activity and retained other characteristic or property, relative 30 at least one other property or characteristic until an antibody, or antigen-binding portion thereof, relative to the parent antibody or antigen-binding portion thereof, for changes in e) evaluating the panel of mutated antibodies, or antigen-binding portions

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In another embodiment, the invention provides a method for improving the

activity of an antibody, or antigen-binding portion thereof, comprising:

- a) providing a parent antibody or antigen-binding portion thereof that was
- obtained by selection in a phage-display system but whose activity cannot be further
- improved by mutagenesis in said phage-display system;

 b) selecting an amino acid residue within a complementarity determining region
- (CDR) for mutation other than H30, H31, H31B, H32, H33, H35, H50, H52, H52A,
- H23' H24' H26' H28' H32' H36' H30' H38' H101' L30, L31, L32, L34, L50, L52, L53,
- L55, L91, L92, L93, L94 and L96;

 c) individually mutating said selected position to at least two other amino acid
- residues to thereby create a panel of mutated antibodies, or antigen-binding portions
- thereof and expression in a non-phage display system;
- d) evaluating the activity and retention of at least one other characteristic or property of the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent entibody or entities property.
- property of the panel of mutated antibodies, of antigen-binding portions thereby identifying an activity enhancing amino acid residue;
- e) repeating steps b) through d) for at least one other CDR position which is neither the position selected under b nor other than H30, H31, H31B, H32, H33, H35, H50, H52, H52A, H53, H54, H56, H58, H96, H97, H98, H101, L30, L31, L32,

F34' F20' F25' F23' F22' F61' F65' F63' F64 and L96;

f) combining, in the parent antibody, or antigen-binding portion thereof, at least two individual activity enhancing amino acid residues shown to have improved activity and not to affect at least one other characteristic or property, to form combination antibodies, or antigen-binding portions thereof; and

g) evaluating the activity and retention of at least one other characteristic or

property of the combination antibodies, or antigen-binding portions thereof with two activity enhancing amino acid residues, relative to the parent antibody or antigen-binding portion thereof, until an antibody, or antigen-binding portion thereof, with an improved activity and at least one other retained characteristic or property, relative to the parent antibody, or antigen-binding portion thereof, is obtained.

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Preferably, the other characteristic or property is selected from 1) preservation of non-crossreactivity with other proteins or human tissues, 2) preservation of epitope recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 heterodimer preventing binding interference from free, soluble p40 and/or 3) to produce an antibody with a close to germline immunoglobulin sequence

Brief Description of the Drawings

Figures 1A-1B show the heavy chain variable region amino acid sequence alignments of a series of human antibodies that bind human IL-12 compared to germline sequences Cos-3/JH3 and Dpl18 Lv1042. Kabat numbering is used to identify amino acid positions. For the Joe 9 wild type, the full sequence is shown. For the other antibodies, only those amino acids positions that differ from Joe 9 wild type are shown. Figures 1C-1D show the light chain variable region amino acid sequence alignments of a series of human antibodies that bind human IL-12. Kabat numbering is

alignments of a series of human antibodies that bind human IL-12. Kabat numbering is used to identify amino acid positions. For the loe 9 wild type, the full sequence is shown. For the other antibodies, only those amino acids positions that differ from loe 9 wild type are shown.

Figures 2A-2E show the CDR positions in the heavy chain of the Y61 antibody that were mutated by site-directed mutagenesis and the respective amino acid

substitutions at each position. The graphs at the right of the figures show the off-rates for the substituted antibodies (black bars) as compared to unmutated Y61 (open bar). Figures 2F-2H show the CDR positions in the light chain of the Y61 antibody

that were mutated by site-directed mutagenesis and the respective amino acid substitutions at each position. The graphs at the right of the figures show the off-rates for the substituted antibodies (black bars) as compared to unmutated Y61 (open bar). Figure 3 demonstrates the *in vivo* efficacy of the human anti-IL-I2 antibody

1695, on plasma neopterin levels in cynomolgus monkeys.

Figure 4 shows a graph of mean arthritic score versus days after immunization of mice with collagen, demonstrating that treatment with C17.15 significantly decreases arthritis-related symptoms as compared to treatment with rat IgG.

Detailed Description of the Invention

In order that the present invention may be more readily understood, certain terms

are first defined.

The term "activity enhancing amino acid residue" includes an amino acid residue

which improves the activity of the antibody. It should be understood that the activity

enhancing amino acid residue may replace an amino acid residue at a contact, hypermutation or preferred selective mutagenesis position and, further, more than one activity enhancing amino acid residue include. an amino acid residue that improves the binding specificity/affinity of an antibody, for example anti-human IL-12 antibody binding to human IL-12. The activity enhancing amino acid residue is also intended to binding to amino acid residue that improves the reclude an amino acid residue that improves the include an amino acid residue that improves the neutralization potency of an antibody,

for example, the human IL-12 antibody which inhibits human IL-12.

The term "antibody" includes an immunoglobulin molecule comprised of four polypeptide chains, two heavy (H) chains and two light (L) chains inter-connected by disulfide bonds. Each heavy chain is comprised of a heavy chain constant region. The heavy chain is comprised of three domains, CH1, CH2 and CH3. Each light chain is comprised of three domains, CH1, CH2 and CH3. Each light and a light chain constant region. The light chain constant region is comprised of one and a light chain constant region. The light chain constant region is comprised of one domain, CL. The VH and VL regions can be further subdivided into regions of hypervariability, termed complementarity determining regions (CDRs), interspersed hypervariability, termed complementarity determining regions (CDRs), interspersed

domain, CL. The VH and VL regions can be further subdivided into regions of hypervariability, termed complementarity determining regions (CDRs), interspersed with regions that are more conserved, termed framework regions (FR). Each VH and VL is composed of three CDRs and four FRs, arranged from amino-terminus to carboxy-terminus in the following order: FRI, CDRI, FR2, CDR2, FR3, CDR3, FR4.

The term "antigen-binding portion" of an antibody (or "antibody portion")

includes fragments of an antibody that retain the ability to specifically bind to an antigen (e.g., hIL-12). It has been shown that the antigen-binding function of an antibody can be performed by fragments of a full-length antibody. Examples of binding fragments
of encompassed within the term "antigen-binding portion" of an antibody include (i) a Fab
fragment, a monovalent fragment consisting of the VL, VH, CL and CH1 domains; (ii) a
F(ab')₂ fragment, a bivalent fragment comprising two Fab fragments linked by a

recombinant DNA techniques, as described herein. Preferred antigen binding portions antibody portions and immunoadhesion molecules can be obtained using standard as papain or pepsin digestion, respectively, of whole antibodies. Moreover, antibodies, fragments, can be prepared from whole antibodies using conventional techniques, such al. (1994) Mol. Immunol. 31:1047-1058). Antibody portions, such as Fab and $F(ab')_2$ polyhistidine tag to make bivalent and biotinylated scFv molecules (Kipriyanov, S.M., et Hybridomas 6:93-101) and use of a cysteine residue, a marker peptide and a C-terminal tetrameric scFv molecule (Kipriyanov, S.M., et al. (1995) Human Antibodies and such immunoadhesion molecules include use of the streptavidin core region to make a antibody or antibody portion with one or more other proteins or peptides. Examples of immunoadhesion molecules, formed by covalent or non-covalent association of the Still further, an antibody or antigen-binding portion thereof may be part of a larger Nail. Acad. Sci. USA 90:6444-6448; Poljak, R.J., et al. (1994) Structure 2:1121-1123). chain and creating two antigen binding sites (see e.g., Holliger, P., et al. (1993) Proc. same chain, thereby forcing the domains to pair with complementary domains of another but using a linker that is too short to allow for pairing between the two domains on the antibodies in which VH and VL domains are expressed on a single polypeptide chain, antibodies, such as diabodies are also encompassed. Diabodies are bivalent, bispecific within the term "antigen-binding portion" of an antibody. Other forms of single chain 85:5879-5883). Such single chain antibodies are also intended to be encompassed (1988) Science 242:423-426; and Huston et al. (1988) Proc. Natl. Acad. Sci. USA pair to form monovalent molecules (known as single chain Fv (scFv); see e.g., Bird et al. that enables them to be made as a single protein chain in which the VL and VH regions by separate genes, they can be joined, using recombinant methods, by a synthetic linker Furthermore, although the two domains of the Fv fragment, VL and VH, are coded for of a VH domain; and (vi) an isolated complementarity determining region (CDR). antibody, (v) a dAb fragment (Ward et al., (1989) Nature 341:544-546), which consists domains; (iv) a Fv fragment consisting of the VL and VH domains of a single arm of an disulfide bridge at the hinge region; (iii) a Fd fragment consisting of the VH and CH1

are complete domains or pairs of complete domains.

The term "backmutation" refers to a process in which some or all of the somatically mutated amino acids of a human antibody are replaced with the corresponding germline residues from a homologous germline antibody of the invention are aligned separately with the germline sequences in the VBASE database to identify the sequences with the highest homology. Differences in the human antibody of the invention are with the highest homology. Differences in the human antibody of the invention are separately with the germline sequences in the human antibody of the invention are

with the highest homology. Differences in the human antibody of the invention are returned to the germline sequence by mutating defined nucleotide positions encoding such different amino acid. The role of each amino acid thus identified as candidate for a backmutation should be investigated for a direct or indirect role in antigen binding and any amino acid found after mutation to affect any desirable characteristic of the human antibody; as an example, activity enhancing amino acids identified by the selective mutagenesis approach will not be subject to backmutation. To minimize the number of amino acids subject to backmutation, those amino acid positions found to be different from the closest germline backmutation those amino acid positions found to be different from the closest germline sequence but identical to the corresponding amino acid in a second germline sequence can remain, provided that the second germline sequence is identical and colinear to the sequence of the human antibody of the invention for at least 10, preferably 12 amino sequence of the human antibody of the invention for at least 10, preferably 12 amino

The phrase "human interleukin 12" (abbreviated herein as hIL-12, or IL-12), as used herein, includes a human cytokine that is secreted primarily by macrophages and dendritic cells. The term includes a heterodimeric protein comprising a 35 kD subunit (p40) which are both linked together with a disulfide bridge.

The heterodimeric protein is referred to as a "p70 subunit". The structure of human IL-12 is described further in, for example, Kobayashi, et al. (1989) J. Exp Med. 170:827-845; Seder, et al. (1993) Proc. Natl. Acad. Sci. 90:10188-10192; Ling, et al. (1995) J. Exp Med. 154:116-127; Podlaski, et al. (1992) Arch. Biochem. Biophys. 294:230-237.

of antibody optimization; preferably, backmutation occurs directly before or affer the selective mutagenesis approach. More preferably, backmutation occurs directly before

acids, on both sides of the amino acid in question. Backmuation may occur at any stage

The term human IL-12 is intended to include recombinant human IL-12 (rh IL-12),

which can be prepared by standard recombinant expression methods.

the selective mutagenesis approach.

The terms "Kabat numbering", "Kabat definitions and "Kabat labeling" are used interchangeably herein. These terms, which are recognized in the art, refer to a system of numbering amino acid residues which are more variable (i.e. hypervariable) than of numbering amino acid residues in the heavy and light chain variable regions of an antibody, or an antigen binding portion thereof (Kabat et al. (1971) Ann. NY Acad, Sci. 190:382-391 and, Kabat, E.A., et al. (1991) Sequences of Proteins of Immunological Interest, Fifth and, Cabat, E.A., et al. (1991) Sequences of Proteins of Immunological Interest, Fifth and, Cabat, E.A., et al. (1991) Sequences of Proteins of Immunological Interest, Fifth and positions 31 to 35 for CDR1, amino acid positions 50 to 65 for CDR2, and amino acid positions 31 to 35 for CDR3. For the light chain variable region, the hypervariable region ranges from amino acid positions 24 to 34 for CDR1, amino acid positions 50 to 65 for CDR2, and amino acid positions 24 to 34 for CDR1, amino acid positions 50 to 65 for CDR2, and amino acid positions 89 to 97 for CDR3.

The Kabat numbering is used herein to indicate the positions of amino acid modifications made in antibodies of the invention. For example, the Y61 anti-IL-12 antibody can be mutated from serine (S) to glutamic acid (E) at position 31 of the heavy chain CDR1 (H31S \rightarrow E), or glycine (G) can be mutated to tyrosine (Y) at position 94 of the light chain CDR3 (L94G \rightarrow Y).

regions corresponding to human germline immunoglobulin sequences as described by

The term "human antibody" includes antibodies having variable and constant

Kabat et al. (See Kabat, et al. (1991) Sequences of Proteins of Immunological Interest, Fifth Edition, U.S. Department of Health and Human Services, NIH Publication No. 91-3242). The human antibodies of the invention may include amino acid residues not encoded by human germline immunoglobulin sequences (e.g., mutations introduced by tandom or site-specific mutagenesis in vitro or by somatic mutation in vivo), for using the "selective mutagenesis approach" described herein. The human antibody can have at least one position replaced with an amino acid residue, e.g., an activity enhancing amino acid residue which is not encoded by the human germline immunoglobulin sequence. The human antibody can have up to twenty positions or replaced with amino acid residues which are not part of the human germline immunoglobulin sequence. In other embodiments, up to ten, up to five, up to three or up to two positions are replaced. In a preferred embodiment, these replacements are within to two positions are replaced. In a preferred embodiment, these replacements are within

the CDR regions as described in detail below. However, the term "human antibody", as used herein, is not intended to include antibodies in which CDR sequences derived from the germline of another mammalian species, such as a mouse, have been graffed onto human framework sequences.

The phrase "recombinant human antibody" includes human antibodies that are

An "isolated antibody" includes an antibody that is substantially free of other 52 mutagenesis approach or backmutation or both. embodiments, however, such recombinant antibodies are the result of selective naturally exist within the human antibody germline repertoire in vivo. In certain while derived from and related to human germline VH and VL sequences, may not sequences of the VH and VL regions of the recombinant antibodies are sequences that, for human Ig sequences is used, in vivo somatic mutagenesis) and thus the amino acid human antibodies are subjected to in vitro mutagenesis (or, when an animal transgenic NIH Publication No. 91-3242). In certain embodiments, however, such recombinant Immunological Interest, Fifth Edition, U.S. Department of Health and Human Services, immunoglobulin sequences (See Kabat, E.A., et al. (1991) Sequences of Proteins of ςı antibodies have variable and constant regions derived from human germline immunoglobulin gene sequences to other DNA sequences. Such recombinant human expressed, created or isolated by any other means that involves splicing of human e.g., Taylor, L.D., et al. (1992) Nucl. Acids Res. 20:6287-6295) or antibodies prepared, from an animal (e.g., a mouse) that is transgenic for human immunoglobulin genes (see human antibody library (described further in Section III, below), antibodies isolated further in Section II, below), antibodies isolated from a recombinant, combinatorial expressed using a recombinant expression vector transfected into a host cell (described prepared, expressed, created or isolated by recombinant means, such as antibodies

antibodies having different antigenic specificities (e.g., an isolated antibody that specifically binds hIL-12 is substantially free of antibodies that specifically binds antigens other than hIL-12). An isolated antibody that specifically binds hIL-12 may bind IL-12 molecules from other species (discussed in further detail below). Moreover, an isolated antibody may be substantially free of other cellular material and/or an isolated antibody may be substantially free of other cellular material and/or

chemicals.

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A "neutralizing antibody" (or an "antibody that neutralized hIL-12 activity") includes an antibody whose binding to hIL-12 results in inhibition of the biological activity of hIL-12. This inhibition of the biological activity of hIL-12 can be assessed by measuring one or more indicators of hIL-12 biological activity. such as inhibition of human phytohemagglutinin blast proliferation in a phytohemagglutinin blast proliferation in a phytohemagglutinin blast proliferation of the phytohemagglutinin blast proliferation of receptor binding in a human IL-12 receptor

proliferation assay (PHA), or inhibition of receptor binding in a human IL-12 receptor binding assay (see Example 3-Interferon-gamma Induction Assay). These indicators of hIL-12 biological activity can be assessed by one or more of several standard in vitro or in vivo assays known in the art (see Example 3).

The term "activity" includes activities such as the binding specificity/affinity of an antibody for an antigen, for example, an anti-hIL-12 antibody that binds to an IL-12 antibody whose binding to hIL-12 inhibits the biological activity of hIL-12, e.g. inhibition of PHA blast proliferation or inhibition of receptor binding in a human IL-12 receptor binding assay (see Example 3).

The phrase "surface plasmon resonance" includes an optical phenomenon that allows for the analysis of real-time biospecific interactions by detection of alterations in protein concentrations within a biosensor matrix, for example using the BIAcore system (Pharmacia Biosensor AB, Uppsala, Sweden and Piscataway, MJ). For further descriptions, see Example 5 and Jönsson, U., et al. (1993) Ann. Biol. Clin. 51:19-26; Jönsson, U., et al. (1991) Biotechniques 11:620-627; Johnsson, B., et al. (1995) J. Mol. Recognit. 8:125-131; and Johnnson, B., et al. (1991) Anal. Biochem. 198:268-277.

The term "K_{off}", as used herein, is intended to refer to the off rate constant for

dissociation of an antibody from the antibody/antigen complex.

The term " K_d ", as used herein, is intended to refer to the dissociation constant of

a particular antibody-antigen interaction.

The phrase "nucleic acid molecule" includes DNA molecules and RNA molecules. A nucleic acid molecule may be single-stranded or double-stranded, but

nucleic acids encoding antibodies or antibody portions (e.g., VH, VL, CDR3) that bind hIL-12 including "isolated antibodies"), includes a nucleic acid molecule in which the

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preferably is double-stranded DNA.

The phrase "isolated nucleic acid molecule", as used herein in reference to

nucleotide sequences encoding the antibody or antibody portion are free of other nucleotide sequences encoding antibodies or antibody portions that bind antigens other than hIL-12, which other sequences may naturally flank the nucleic acid in human genomic DNA. Thus, for example, an isolated nucleic acid of the invention encoding a VH region of an anti-IL-12 antibody contains no other sequences encoding other VH regions that bind antigens other than IL-12. The phrase "isolated nucleic acid molecule" is also intended to include sequences encoding bivalent, bispecific antibodies, such as diabodies in which VH and VL regions contain no other sequences other than the sequences of the diabody.

vectors (e.g., replication defective retroviruses, adenoviruses and adeno-associated the invention is intended to include such other forms of expression vectors, such as viral interchangeably as the plasmid is the most commonly used form of vector. However, plasmids. In the present specification, "plasmid" and "vector" may be used expression vectors of utility in recombinant DNA techniques are often in the form of "recombinant expression vectors" (or simply, "expression vectors"). In general, to which they are operatively linked. Such vectors are referred to herein as host genome. Moreover, certain vectors are capable of directing the expression of genes host cell upon introduction into the host cell, and thereby are replicated along with the vectors (e.g., non-episomal mammalian vectors) can be integrated into the genome of a vectors having a bacterial origin of replication and episomal mammalian vectors). Other autonomous replication in a host cell into which they are introduced (e.g., bacterial DNA segments may be ligated into the viral genome. Certain vectors are capable of segments may be ligated. Another type of vector is a viral vector, wherein additional which refers to a circular double stranded DNA loop into which additional DNA another nucleic acid to which it has been linked. One type of vector is a "plasmid", The term "vector" includes a nucleic acid molecule capable of transporting

The phrase "recombinant host cell" (or simply "host cell") includes a cell into which a recombinant expression vector has been introduced. It should be understood that such terms are intended to refer not only to the particular subject cell but to the progeny of such a cell. Because certain modifications may occur in succeeding generations due to either mutation or environmental influences, such progeny may not,

viruses), which serve equivalent functions.

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in fact, be identical to the parent cell, but are still included within the scope of the term "host cell" as used herein.

The term "modifying", as used herein, is intended to refer to changing one or

more amino acids in the antibodies or antigen-binding portions thereof. The change can be produced by adding, substituting or deleting an amino acid at one or more positions. The change can be produced using known techniques, such as PCR mutagenesis.

The phrase "contact position" includes an amino acid position of in the CDRI, or CDR2 or CDR3 of the heavy chain variable region or the light chain variable region of the an amino acid that contacts antigen in one of the twenty-six known antibody-antigen structures. If a CDR amino acid in any of the 26 known solved structures of antibody-antigen complexes contacts the antigen, then that amino acid can be considered to occupy a contact position. Contact positions have a higher probability of being occupied by an amino acid which contact antigen than noncontact positions. Preferably a contact position is a CDR position which contains an amino acid that contacts position is a CDR position which contains an amino acid that preferably a contact position is a CDR position which contains an amino acid that

The term "hypermutation position" includes an amino acid residue that occupies position in the CDR1, CDR2 or CDR3 region of the heavy chain variable region or the probability for somatic hypermutation during in vivo affinity maturation of the antibody. "High frequency or probability for somatic hypermutation during in vivo affinity maturation of the antibody. "High frequency or probability for somatic hypermutation" includes frequencies or "High frequency or probability for somatic hypermutation during in vivo affinity maturation" includes frequencies or hypermutation during in vivo affinity maturation of the antibody. It should be

contacts antigen in greater than 8 of the 25 structures (>32%).

understood that all ranges within this stated range are also intended to be part of this invention, e.g., 5 to about 30%, e.g., 5 to about 15%, e.g., 15 to about 30%.

The term "preferred selective mutagenesis position" includes an amino acid

residue that occupies a position in the CDRI, CDR2 or CDR3 region of the heavy chain variable region or the light chain variable region which can be considered to be both a contact and a hypermutation position.

germline genes, human antibodies isolated from human B-cells. Preferably, the selective derived from any source including phage display, transgenic animals with human IgG 30 selective mutagenesis approach can be used for the optimization of any antibody antibody, and/or neutralization potency of the antibody. It should be understood that the is determined. Activity is measured as a change in the binding specificity/affinity of the other amino acid residues, and the effect of the mutation on the activity of the antibody H2, L1, H1 and L2. The selected amino acid residue is mutated, e.g., to at least two 52 for selective mutations with the order of preference for targeting as follows: H3, L3, wise targeted approach", individual amino acid residues in particular CDRs are targeted L2 and H1), the groups being listed in order of preference for targeting. In the "CDRincluding groups I (including L3 and H3), II (including H2 and L1) and III (including individual amino acid residues in particular groups are targeted for selective mutations 50 approach" or "CDR-wise targeted approach". In the "Group-wise targeted approach", chain variable region of an antibody in a targeted manner, e.g., a "Group-wise targeted CDR3 of the heavy chain variable region or the CDR1, CDR2 or CDR3 of the light preferentially mutating selected individual amino acid residues in the CDR1, CDR2 or approach". The language "targeted approach" is intended to include a method of contact position. In an embodiment, the selective mutagenesis approach is a "targeted variable region. It should be understood that a hypermutation position can also be a Individual amino acids are selected based on their position in the light or heavy chain preferred selective mutagenesis positions, contact positions., or hypermutation positions. L1, L2, and L3, respectively) of an antibody. Amino acid residues may be selected from the CDR1, CDR2 or CDR3 of the light chain variable region (hereinafter referred to as CDR3 of the heavy chain variable region (hereinafter H1, H2, and H3, respectively), or preferentially mutating selected individual amino acid residues in the CDRI, CDR2 or embodiment, the selective mutagenesis approach is intended to provide a method of mutation at a position selected using a selective mutagenesis approach. In another position. A "selectively mutated" human antibody is an antibody which contains a least one preferred selective mutagenesis position, hypermutation, and/or contact activity of an antibody by selecting and individually mutating CDR amino acids at at The phrase "selective mutagenesis approach" includes a method of improving the

mutagenesis approach is used on antibodies which can not be optimized further using

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phage display technology. It should be understood that antibodies from any source including phage display, transgenic animals with human IgG germline genes, human antibodies isolated from human B-cells can be subject to backmutation prior to or after the selective mutagenesis approach.

The term "activity enhancing amino acid residue" includes an amino acid residue

which improves the activity of the antibody. It should be understood that the activity enhancing amino acid residue may replace an amino acid residue at a preferred selective mutagenesis position, contact position, or a hypermutation position and, further, more than one activity enhancing amino acid residue can be present within one or more CDRs.

An activity enchancing amino acid residue include, an amino acid residue that improves the binding specificity/affinity of an antibody, for example anti-human IL-12 antibody binding to human IL-12. The activity enhancing amino acid residue is also intended to include an amino acid residue that improves the neutralization potency of an antibody, for example, the human IL-12 antibody which inhibits human IL-12.

Various aspects of the invention are described in further detail in the following

This invention provides isolated human antibodies, or antigen-binding portions

subsections.

I. Human Antibodies that Bind Human IL-12

alignments of Figures 1A-D).

thereof, that bind to human IL-12. Preferably, the human antibodies of the invention are recombinant, neutralizing human anti-hIL-12 antibodies. Antibodies of the invention that bind to human IL-12 can be selected, for example, by screening one or more human VL and VH cDNA libraries with hIL-12, such as by phage display techniques as described in Example 1. Screening of human VL and VH cDNA libraries initially identified a series of anti-IL-12 antibodies of which one antibody, referred to herein as low affinity human IL-12 antibody (e.g., a K_{off} of about 0.1 sec⁻¹), yet is useful for low affinity human IL-12 antibody (e.g., a K_{off} of about 0.1 sec⁻¹), yet is useful for specifically binding and detecting hIL-12. The affinity of the Joe 9 antibody was improved by conducting mutagenesis of the heavy and light chain CDRs, producing a further mutated, leading to numerous additional anti-hIL-12 antibodies with increased further mutated, leading to numerous additional anti-hIL-12 antibodies with increased affinity for hIL-12 (see Example 1, Table 2 (see Appendix A) and the sequence

light chain CDR2 of Y61, and a Gly to Tyr substitution at position 94 of the light chain referred to herein as J695, resulted from a Gly to Tyr substitution at position 50 of the Figures 2A-2H. A preferred recombinant neutralizing antibody of the invention, substitutions at selected positions in the heavy and light chain CDRs is shown in mutagenesis position, contact and/or a hypermutation position. A summary of the mutagenesis approach) based on the amino acid residue occupying a preserved selective CDRs. Amino acids residues of Y61 were selected for site-specific mutation (selective individually mutating specific amino acids residues within the heavy and light chain sec-1). The Y61 anti-hIL-12 antibody was selected for further affinity maturation by demonstrated a significant improvement in binding affinity (e.g., a K_{off} of about 2 x 10^{-4} Of these antibodies, the human anti-hlL-12 antibody referred to herein as Y61

for the CDR3, CDR2, and CDR1, on the lineage from loe 9 to 1695. Moreover, the regions of antibodies of the invention that bind hIL-I2, as well as consensus sequences identification of consensus sequences for preferred heavy and light chain variable J695, are shown in Figures 1A-1D. These sequence alignments allowed for the a panel of anti-IL-12 antibodies of the invention, on the lineage from loe 9 wild type to Amino acid sequence alignments of the heavy and light chain variable regions of

CDK3 of Y61.

the attached Sequence Listing, are summarized below. of the invention (including consensus sequences) as identified by sequence identifiers in that retain good hIL-12 binding characteristics. Preferred CDR, VH and VL sequences lineage from Y61 to 1695 that encompasses sequences with modifications from Y61 yet well as consensus sequences for the CDR3, CDR2, and CDR1 that bind hIL-12 on the of consensus sequences for heavy and light chain variable regions that bind hlL-12, as Y61 mutagenesis analysis summarized in Figures 2A-2H allowed for the identification

Ŏ-(S\L)-X-(D\E)-(S\K\K)-(S\C\X)-	CDK F3	Consensus	7
		369r of 6 9or	
(H/S) -G-S-(H/X) -D-(N/T/X)	СDК НЗ	Consensus	τ
			:ОИ
		CHYIN	ID
геблеисе	KECION	YOOHITMA	SEŎ

-	91	7	

	·		
S-G-G-K-S-N-I-G-S-N-L-A-K	CDK PJ	٨٤٦	22
F-T-F-S-Z-Y-H	СDК НТ	Т9Х	TZ
G-N-D-Ŏ-K-Þ-2	CDK PS	τ9λ	20
E-I-K-X-D-G-Z-M-K-X-X-Y-D-Z-A-K-G	CDK HS	τ9λ	6 T
Ŏ-8-X-D-K-G-T-H-P-A-L-L	срк гз	T9X	18
н-д-к-р-и	СDК НЗ	τ9λ	L٦
sedneuce listing)		sear oa tay	
(Įnjj Ar sedneuce; see	ΛΓ	Consensus	9τ
sedneuce listing)		263t cd 18Y	
(fnll VH sequence; see	НΛ	Consensus	ST
(N/W/I) - (L/X/D/H/K/b) - A-K		Tel to Jess	
2-G-G-Б-2-И-I-G-(2\C\K\N\D\L)-	CDK L1	Consensus	ÐΤ
H-M-(S/N/A/M/Ð)		Xet to Jess	
- (T-T-T-T-T-T-T-T-T-T-T-T-T-T-T-T-T-T-T	CDK HJ	Consensus	ΣŢ
		269C 03 L9Y	
(G/Y/S/T/N/Q) -N-D-Q-R-P-S	CDK PS	Consensus	75
2-(X99)-K-(X\E)-X-Y-D-2-A-K-G		269£ 63 £9X	
(E/L/X)-I-(K/Y)-X-(D/S/E/Y)-(G/K)-	CDK HS	Consensus	TT
(G/D/Ŏ/r/Ł/K/H/N/X)-T-H-P-A-L-L		569£ 07 £9X	
- (kak) - (V/G) -Y-2-Q	CDK P3	Consensus	01
(D/S) - (N/K/Y/L/S/E/M/H)		269r of 19Y	
H-(G\\\C\H)-(S\\L)-(H\\\\\\L)-	CDK H3	Consensus	6
sequence listing)		269r of 6 90r	
(fn]] Ar sedneuce; see	ΛΓ	Consensus	8
sednence listing)		10e 9 to 1695	
(fn]] VH sequence; see	НΛ	Consensus	L
(B/A) - (A/B) - (T/D) - V - (K/H)		269r of 6 9or	
$(2\L) - G - (G\Z) - (B\Z) - 2 - N - I - (G\A) -$	CDK TJ	Consensus	9
,		269r of 6 9or	
E-T-E-S-(S/E)-X-G-M-H	CDK HJ	Consensus	<u>S</u>
		269r oj 6 90r	
(G\X) -N- (D\R) - (Ō\N) -K- <u>b-</u> R	CDK PS	Consensus	ъ
		269L 01 6 90L	
E-I-B-Y-D-G-S-N-K-Y-Y-D-S-V-K-G	CDK HS	Consensus	3
(J/M/T/\V) - (J/T/W/S/A) - (J/A/T/S)			
$(\Gamma \setminus E \setminus T \setminus S) - (E \setminus S \setminus T \setminus M \setminus H) - (G \setminus D) -$		269r of 6 9or	

sequence listing)			
([n]] Nr sedneuce; see	ΛΓ	569L	32
sedneuce listing)			
(fnll VH sequence; see	НΛ	569C	3.1
Z-G-Z-K-Z-N-I-G-Z-N-L-A-K	СDК ГЈ	569C	30
F-T-F-S-Z-Y-H	CDK HJ	269L	52
X-N-D-Ğ-K-D-Z	CDK PS	269r	28
E-I-K-X-D-G-2-M-K-X-X-Y-D-2-A-K-G	CDK HS	269r	LZ
Ŏ-а-х-D-К-х- Т -H-Ъ-У-Г-Г	CDK F3	569r	97
И-С-В-Б-И	СDИ НЗ	269L	52
sedneuce listing)			
(inll VL sequence; see	ΛΓ	τ9λ	5₫
sednence listing)	•		
(£n]] AH sedneuce: see	НΛ	Τ9.Κ	23

Antibodies produced from affinity maturation of loe 9 wild type were

Accordingly, in one aspect, the invention provides an isolated human antibody, 1x10-11 M, more preferably about 1x10-10 M to 1x10-11 M or less. antibodies were produced having an IC50 value in the range of about 1x10-6 M to about phytohemagglutinin (PHA) blast proliferation, as described in Example 3. A series of I x 10-5 s-1 or less. Antibodies were also characterized in vitro for their ability to inhibit of about 0.1 s⁻¹ to about 1 x 10⁻⁵ s⁻¹, and more preferably a K_{off} of about 1 x 10⁻⁴ s⁻¹ to and K_{off} rate. A series of antibodies were produced having a K_{off} rate within the range functionally characterized by surface plasmon resonance analysis to determine the $K_{
m d}$

assay with an IC₅₀ of 1 x 10-7 M or less. In more preferred embodiments, the isolated I x 10-2 s-1 or less, or inhibits phytohemagglutinin blast proliferation in an in vitro PHA binding portion thereof, dissociates from human IL-12 with a Koff rate constant of M or less. In preferred embodiments, the isolated human IL-12 antibody, or an antigenvitro phytohemagglutinin blast proliferation assay (PHA assay) with an IC₅₀ of 1 x 10-6 plasmon resonance, or which inhibits phytohemagglutinin blast proliferation in an in human IL-12 with a K_{off} rate constant of $0.1~\text{s}^{-1}$ or less, as determined by surface or antigen-binding portion thereof, that binds to human IL-12 and dissociates from

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IL-12 with a K_{off} rate constant of 1 x 10⁻³ s⁻¹ or less, or inhibits phytohemagglutinin human IL-12 antibody, or an antigen-binding portion thereof, dissociates from human

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blast proliferation in an *in vitro* PHA assay with an IC₅₀ of 1 x 10-8 M or less. In more preferred embodiments, the isolated human IL-12 antibody, or an antigen-binding portion thereof, dissociates from human IL-12 with a K_{off} rate constant of 1 x 10-4 s⁻¹ or less, or inhibits phytohemagglutinin blast proliferation in an *in vitro* PHA assay with an antibody, or an antigen-binding portion thereof, dissociates from human IL-12 with a more preferred embodiments, the isolated human IL-12 with a proliferation in an *in vitro* PHA assay with an IC₅₀ of 1 x 10-10 M or less. In even more preferred embodiments, the isolated human IL-12 antibody, or an antigen-binding portion thereof, dissociates from human IL-12 antibody, or an antigen-binding portion thereof, dissociates from human IL-12 with a K_{off} rate constant of 1 x 10-5 s⁻¹ or portion thereof, dissociates from human IL-12 with a K_{off} rate constant of 1 x 10-5 s⁻¹ or portion thereof, dissociates from human IL-12 with a K_{off} rate constant of 1 x 10-5 s⁻¹ or less, or inhibits phytohemagglutinin blast proliferation in an *in vitro* PHA assay with an essay or inhibits phytohemagglutinin blast proliferation in an *in vitro* PHA assay with an

 IC_{50} of 1×10^{-11} M or less.

surface plasmon resonance (see Example 5). Generally, surface plasmon resonance analysis measures real-time binding interactions between ligand (recombinant human IL-12 immobilized on a biosensor matrix) and analyte (antibodies in solution) by surface plasmon resonance (SPR) using the BIAcore system (Pharmacia Biosensor, Piscataway, AJ). Surface plasmon analysis can also be performed by immobilizing the analyte (antibodies on a biosensor matrix) and presenting the ligand (recombinant IL-12 in solution). Meutralization activity of IL-12 antibodies, or antigen binding portions aclution). Acutalization activity of IL-12 antibodies, or antigen binding portions between the analyte and a biosensor matrix.

The dissociation rate constant (K_{off}) of an IL-12 antibody can be determined by

It is well known in the art that antibody heavy and light chain CDRs play an

important role in the binding specificity/affinity of an antibody for an antigen.

CDRs of Joe 9, as well as other antibodies having CDRs that have been modified to improve the binding specificity/affinity of the antibody. As demonstrated in Example 1, a series of modifications to the light and heavy chain CDRs results in affinity maturation of human anti-hIL-12 antibodies. The heavy and light chain variable region amino acid sequence alignments of a series of human antibodies ranging from Joe 9 wild type to J695 that bind human IL-12 is shown in Figures 1A-1D. Consensus sequence type to J695 that bind human IL-12 is shown in Figures 1A-1D. Consensus sequence motifs for the CDRs of antibodies can be determined from the sequence alignment (as

summatized in the table above). For example, a consensus motif for the VH CDR3 of the lineage from loe 9 to 1695 comprises the amino acid sequence: (H/S)-G-S-(H/Y)-D-(N/T/Y) (SEQ ID NO: 1), which encompasses amino acids from position 95 to 102 of the consensus HCVR shown in SEQ ID NO: 7. A consensus motif for the VL CDR3 comprises the amino acid sequence: Q-(S/T)-Y-(D/E)-(S/R/K)-(S/G/Y)-(L/F/T/S)-(R/S/T/W/H)-(G/P)-(S/T/A/L)-(R/S/M/T/L-V/I/T/M/L) (SEQ ID NO: 2), which encompasses amino acids from position 89 to 97 of the consensus LCVR shown in SEQ encompasses amino acids from position 89 to 97 of the consensus LCVR shown in SEQ

Accordingly, in another aspect, the invention provides an isolated human

antibody, or an antigen-binding portion thereof, which has the following characteristics:

a) inhibits phytohemagglutinin blast proliferation in an in vitro PHA assay

ID NO: 8.

with an IC $_{50}$ of 1 x 10-6 M or less; b) has a heavy chain CDR3 comprising the amino acid sequence of SEQ ID

NO: 1; and

NO: 2.

NO: 2.

In a preferred embodiment, the antibody further comprises a VH CDR2 comprising the amino acid sequence: F-I-R-Y-D-G-S-N-K-Y-A-D-S-V-K-G (SEQ ID NO: 3) (which encompasses amino acid sequence SEQ ID NO: 7) and further comprises a VL CDR2 comprising the amino acid sequence: (G/Y)-N-(D/S)-(Q/N)-R-P-S (SEQ ID NO: 4) (which encompasses amino acid sequence SEQ ID NO: 6) of the consensus LCVR comprising the amino acid sequence SEQ ID NO: 8).

comprising the amino acid sequence: F-T-F-S-(S/E)-Y-G-M-H (SEQ ID NO: 5) (which encompasses amino acids from position 27 to 35 of the consensus HCVR comprising the amino acid sequence: (S/T)-G-(G/S)-(R/S)-S-N-I-(G/V)-(S/A)-(N/G/Y)-(T/D)-Y-(K/H) (SEQ ID NO: 6) (which encompasses amino acids from position 24 to 34 of the consensus LCVR comprising the amino acid sequence SEQ ID NO: 8).

In another preferred embodiment, the antibody further comprises a VH CDR1

HCVR comprising the amino acid sequence of SEQ ID NO: 7 and a LCVR comprising In yet another preferred embodiment, the antibody of the invention comprises a

Additional consensus motifs can be determined based on the mutational analysis the amino acid sequence of SEQ ID NO: 8.

substitution with a variety of different amino acid residues. Thus, based on the reduce the Koff rate of the antibody, indicating that is position is amenable to position 30 in CDR H1 with twelve different amino acid residues did not significantly the hIL-12 binding properties of the antibody. For example, individual substitutions at light chain CDRs of Y61 were amenable to substitution without significantly impairing demonstrated by the graphs shown in Figures 2A-2H, certain residues of the heavy and performed on Y61 that led to the J695 antibody (summarized in Figures 2A-2H). As

and 12, respectively, and consensus motifs for the heavy and light chain CDRIs are consensus motifs for the heavy and light chain CDR2s are shown in SEQ ID NOs: 11 the heavy and light chain CDR3s are shown in SEQ ID NOs: 9 and 10, respectively, other amino acid residues) consensus motifs were determined. The consensus motifs for mutational analysis (i.e., positions within Y61 that were amenable to substitution by

regions are shown in SEQ ID NOs: 15 and 16, respectively. shown in SEQ ID NOs: 13 and 14, respectively. Consensus motifs for the VH and VL

an antigen-binding portion thereof, which has the following characteristics: Accordingly, in one aspect, the invention features an isolated human antibody, or

has a heavy chain CDR3 comprising the amino acid sequence of SEQ ID (q with an IC_{50} of 1 x 10-9 M or less;

inhibits phytohemagglutinin blast proliferation in an in vitro PHA assay

has a light chain CDR3 comprising the amino acid sequence of SEQ ID (c) 52 NO: 9; and

CDR2 comprising the amino acid sequence of SEQ ID NO: 12. comprising the amino acid sequence of SEQ ID NO: 11 and further comprises a VL In a preferred embodiment, the antibody further comprises a VH CDR2 NO: 10.

CDR1 comprising the amino acid sequence of SEQ ID NO: 14. comprising the amino acid sequence of SEQ ID NO: 13 and further comprises a VL In another preferred embodiment, the antibody further comprises a VH CDR1

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In yet another preferred embodiment, the antibody of the invention comprises a HCVR comprising the amino acid sequence of SEQ ID NO: 15 and a LCVR comprising the amino acid sequence of SEQ ID NO: 16.

A preferred antibody of the invention, the human anti-hIL-12 antibody Y61, was

produced by affinity maturation of Joe 9 wild type by PCR mutagenesis of the CDR3 (as described in Example 1). Y61 had an improved specificity/binding affinity determined by surface plasmon resonance and by in vitro neutralization assays. The heavy and light chain CDR3s of Y61 are shown in SEQ ID NOs: 17 and 18, respectively, the heavy and light chain CDR2s of Y61 are shown in SEQ ID NOs: 19 and 20, respectively, and the

10 heavy and light chain CDR1s of Y61 are shown in SEQ ID NOs: 21 and 22, respectively. The VH of Y61 has the amino acid sequence of SEQ ID NO: 24 (these sequences are also shown in Figures 1A-1D, aligned with Joe9).

Accordingly, in another aspect, the invention features an isolated human

antibody, or an antigen-binding portion thereof, which

- a) inhibits phytohemagglutinin blast proliferation in an in vitro PHA assay
- with an IC_{50} of 1 x 10-9 M or less; by has a heavy chain CDR3 comprising the amino acid sequence of SEQ ID
- NO: 17; and

 c) has a light chain CDR3 comprising the amino acid sequence of SEQ ID

In a preferred embodiment, the isolated human antibody, or an antigen-binding portion thereof, has a heavy chain CDR2 comprising the amino acid sequence of SEQ ID MO: 19 and a light chain CDR2 comprising the amino acid sequence of SEQ ID MO: 20.

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NO: 18.

In another preferred embodiment, the isolated human antibody, or an antigen-binding portion thereof has a heavy chain CDR1 comprising the amino acid sequence of SEQ ID NO: 21 and a light chain CDR1 comprising the amino acid sequence of SEQ ID NO: 22.

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In yet another preferred embodiment, the isolated human antibody, or an antigenbinding portion thereof, comprising a the heavy chain variable region comprising the amino acid sequence of SEQ ID NO: 23, and a light chain variable region comprising the amino acid sequence of SEQ ID NO: 24.

In certain embodiments, the full length antibody comprises a heavy chain

constant region, such as IgG1, IgG2, IgG4, IgM, IgA and IgE constant regions, and any allotypic variant therein as described in Kabat (, Kabat, E.A., et al. (1991)

Sequences of Proteins of Immunological Interest, Fifth Edition, U.S. Department of

Health and Human Services, MIH Publication No. 91-3242). Preferably, the antibody

one and Fab fragment, an IgG1 heavy chain constant region. Alternatively, the antibody portion can be an Fab fragment, an F(ab'2) fragment or a single chain Fv

fragment.

Modifications of individual residues of Y61 led to the production of a panel of antibodies shown in Figures 2A-2H. The specificity/binding affinity of each antibody was determined by surface plasmon resonance and/or by in vitro neutralization assays. Accordingly, in another aspect, the invention features an isolated human

antibody, or an antigen-binding portion thereof, which

a) inhibits phytohemagglutinin blast proliferation in an in vitro PHA assay

- with an IC_{50} of 1 x 10-9 M or less; 20 b) has a heavy chain CDR3 comprising the amino acid sequence selected
- from the group consisting of SEQ ID NO: 404-SEQ ID NO: 469; and c) has a light chain CDR3 comprising the amino acid sequence selected
- from the group consisting of SEQ ID NO: 534-SEQ ID NO: 579.

 In preferred embodiment, the isolated human antibody, or an antigen-binding portion thereof, has a heavy chain CDR2 comprising the amino acid sequence selected from the group consisting of SEQ ID NO: 335-SEQ ID NO: 403; and a light chain CDR2 comprising the amino acid sequence selected from the group consisting of SEQ ID NO:
- In another preferred embodiment, the isolated human antibody, or an antigenbinding portion thereof, has a heavy chain CDRI comprising the amino acid sequence selected from the group consisting of SEQ ID NO: 288-SEQ ID NO: 334; and a light

200-SEÓ ID NO: 233.

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2EÓ ID NO: 410-2EÓ ID NO: 202' chain CDR1 comprising the amino acid sequence selected from the group consisting of

amino acid sequence of SEQ ID NO: 23, and a light chain variable region comprising binding portion thereof, comprising a the heavy chain variable region comprising the In yet another preferred embodiment, the isolated human antibody, or an antigen-

the amino acid sequence of SEQ ID NO: 24.

heavy chain constant region is an IgO1 heavy chain constant region. Alternatively, the Health and Human Services, NIH Publication No. 91-3242). Preferably, the antibody Sequences of Proteins of Immunological Interest. Fifth Edition, U.S. Department of any allotypic variant therein as described in Kabat (, Kabat, E.A., et al. (1991) constant region such as IgG1, IgG2, IgG4, IgM, IgA and IgE constant regions and In certain embodiments, the full length antibody comprising a heavy chain

antibody portion can be a Fab fragment, an F(ab)2) fragment or a single chain Fv

light chain CDR2s of J695 are shown in SEQ ID NOs: 27 and 28, respectively, and the chain CDR3s of 1695 are shown in SEQ ID MOs: 25 and 26, respectively, the heavy and Gly to Tyr substitution at position 94 of the light chain CDR3. The heavy and light Y61 by a Gly to Tyr substitution in Y61 at position 50 of the light chain CDR2 and by a acids residues of antibody Y61 (see Example 2 and section III below). 1695 differs from J695, was produced by site-directed mutagenesis of contact and hypermutation amino A particularly preferred recombinant, neutralizing antibody of the invention,

 $\rm VL$ of 1695 has the amino acid sequence of SEQ ID NO: 32 (these sequences are also respectively. The VH of J695 has the amino acid sequence of SEQ ID NO: 31 and the heavy and light chain CDR1s of 1695 are shown in SEQ ID MOs: 29 and 30,

Accordingly, in another aspect, the invention features an isolated human shown in Figures 1A-1D, aligned with Joe9).

inhibits phytohemagglutinin blast proliferation in an in vitro PHA assay g) antibody, or an antigen-binding portion thereof, which

30 with an IC_{50} of 1 x 10-9 M or less;

has a heavy chain CDR3 comprising the amino acid sequence of SEQ ID (q

NO: 25; and

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c) has a light chain CDR3 comprising the amino acid sequence of SEQ ID

NO: 26.

In preferred embodiment, the isolated human antibody, or an antigen-binding portion thereof, has a heavy chain CDR2 comprising the amino acid sequence of SEQ ID.

ID NO: 27, and a light chain CDR2 comprising the amino acid sequence of SEQ ID.

NO: 28.

In another preferred embodiment, the isolated human antibody, or an antigen-binding portion thereof, has a heavy chain CDR1 comprising the amino acid sequence of SEQ ID NO: 29, and a light chain CDR1 comprising the amino acid sequence of

2EÓ ID NO: 30°.

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In yet another preferred embodiment, the isolated human antibody, or an antigen-binding portion thereof, has a heavy chain variable region comprising the amino seid sequence of SEQ ID NO: 31, and a light chain variable region comprising the amino acid sequence of SEQ ID NO: 32.

constant region, such as IgG1, IgG2, IgG4, IgM, IgA and IgE constant regions and any allotypic variant therein as described in Kabat (, Kabat, E.A., et al. (1991)

Sequences of Proteins of Immunological Interest, Fifth Edition, U.S. Department of Health and Human Services, VIH Publication No. 91-3242). Preferably, the antibody portion can be an Fab fragment, an F(ab'2) fragment or a single chain Fv fragment.

methods of modification can be performed using standard molecular biology techniques, such as by PCR mutagenesis, targeting individual contact or hypermutation amino acid residues in the light chain and/or heavy chain CDRs-, followed by kinetic and functional analysis of the modified antibodies as described herein (e.g., neutralization assays described in Example 3, and by BIA core analysis, as described in Example 5).

and CDR1 of antibodies on the lineage from loe 9 to 1695, or from the lineage Y61 to

Additional mutations in the preferred consensus sequences for CDR3, CDR2,

Accordingly, in another aspect the invention features an isolated human

antibody, or an antigen-binding portion thereof, which

a) inhibits phytohemagglutinin blast proliferation in an in vitro PHA assay with an IC $_{50}$ of 1 \times 10-6 M or less;

comprises a heavy chain CDR3 comprising the amino acid sequence of

- SEQ ID MO: 1, a heavy chain CDR1 comprising the amino acid sequence of SEQ ID MO: 5, and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID MO: 5, and a mutant thereof having one or more amino acid substitutions at a preferred selective mutagenesis position or a hypermutation position, wherein said mutant has a k_{off} rate no more than 10-fold higher than the antibody comprising a heavy chain CDR3 comprising the amino acid sequence of SEQ ID MO: 3, and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID MO: 3, and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID MO: 3, and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID MO: 5, and
- SEQ ID NO: 2, a light chain CDR2 comprising the amino acid sequence of SEQ ID NO: 6, or a mutant thereof having one or more amino acid substitutions at a preferred selective mutagenesis position or a hypermutation position, wherein said mutant has a k_{off} rate no more than 10-fold higher than the antibody comprising a light chain CDR3 comprising the amino acid sequence of SEQ ID NO: 2, a light chain CDR2 comprising the amino

comprises a light chain CDR3 comprising the amino acid sequence of

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- acid sequence of SEQ ID NO: 4, and a light chain CDR1 comprising the amino acid sequence of SEQ ID NO: 6.

 In another aspect the invention features an isolated human antibody, or an
- antigen-binding portion thereof, which
 25 a) inhibits phytohemagglutinin blast proliferation in an in vitro PHA assay
- with an IC₅₀ of 1 x 10-9 M or less;

 b) comprises a heavy chain CDR3 comprising the amino acid sequence of SEQ ID MO: 9, a heavy chain CDR1 comprising the amino acid sequence of SEQ ID MO: 11 and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID MO: 13, or a mutant thereof having one or more amino acid substitutions at a preferred
- selective mutagenesis position, contact position or a hypermutation position, wherein said mutant has a $k_{\rm off}$ rate no more than 10-fold higher than the antibody comprising a

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heavy chain CDR3 comprising the amino acid sequence of SEQ ID NO: 11, and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID NO: 13; and CDR1 comprising the amino acid sequence of SEQ ID NO: 13; and

comprises a light chain CDR3 comprising the amino acid sequence of

SEQ ID NO: 10, a light chain CDR1 comprising the amino acid sequence of SEQ ID NO: 12, and a light chain CDR1 comprising the amino acid sequence of SEQ ID NO: 14, or a mutant thereof having one or more amino acid substitutions at a preferred said mutant has a k_{off} rate no more than 10-fold higher than the antibody comprising a light chain CDR3 comprising the amino acid sequence of SEQ ID NO: 10, a light chain CDR2 comprising the amino acid sequence of SEQ ID NO: 12, and a light chain CDR1

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comprising the amino acid sequence of SEQ ID NO: 14.

An ordinarily skilled artisan will also appreciate that additional mutations to the CDR regions of an antibody of the invention, for example in Y61 or in J695, can be

made to provide additional anti-IL-12 antibodies of the invention. Such methods of modification can be performed using standard molecular biology techniques, as described above. The functional and kinetic analysis of the modified antibodies can be performed as described in Example 3 and Example 5, respectively. Modifications of individual residues of Y61 that led to the identification of J695 are shown in Figures 2A-

O ZH and are described in Example 2.

Accordingly, in another aspect the invention features an isolated human antibody, or an antigen-binding portion thereof, which

antibody, or an antigen-binding portion detect, which an in vitro PHA assay with an IC_{50} of 1 x 10^{-9} M or less;

SEQ ID NO: 17, a heavy chain CDR2 comprising the amino acid sequence of SEQ ID NO: 17, a heavy chain CDR2 comprising the amino acid sequence of SEQ ID NO: 19 and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID NO: 21, or a mutant thereof having one or more amino acid substitutions at a preferred selective mutagenesis position or a hypermutation position, wherein said mutant has a koff rate no more than 10-fold higher than the antibody comprising a heavy chain CDR3 comprising the amino acid sequence of SEQ ID NO: 17, a heavy chain CDR3 comprising the amino acid sequence of SEQ ID NO: 17, a heavy chain CDR2

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comprising the amino acid sequence of SEQ ID NO: 21; and comprising the amino acid sequence of SEQ ID NO: 19, and a heavy chain CDR1

- NO: 20, and a light chain CDR1 comprising the amino acid sequence of SEQ ID NO: SEQ ID NO: 18. a light chain CDR2 comprising the amino acid sequence of SEQ ID comprises a light chain CDR3 comprising the amino acid sequence of
- the amino acid sequence of SEQ ID NO: 20, and a light chain CDR1 comprising the comprising the amino acid sequence of SEQ ID NO: 18, a light chain CDR2 comprising koff rate no more than 10-fold higher than the antibody comprising a light chain CDR3 selective mutagenesis position or a hypermutation position, wherein said mutant has a 22, or a mutant thereof having one or more amino acid substitutions at a preferred
- In another aspect the invention features an isolated human antibody, or an amino acid sequence of SEQ ID NO: 22.
- inhibits phytohemagglutinin blast proliferation in an in vitro PHA assay antigen-binding portion thereof, which
- comprising the amino acid sequence of SEQ ID NO: 27, and a heavy chain CDR 1 comprising the amino acid sequence of SEQ ID NO: 25, a heavy chain CDR2 koff rate no more than 10-fold higher than the antibody comprising a heavy chain CDR3 selective mutagenesis position or a hypermutation position, wherein said mutant has a 29, or a mutant thereof having one or more amino acid substitutions at a preferred NO: 27 and a heavy chain CDRI comprising the amino acid sequence of SEQ ID NO: SEQ ID NO: 25, a heavy chain CDR2 comprising the amino acid sequence of SEQ ID comprises a heavy chain CDR3 comprising the amino acid sequence of with an IC_{50} of 1 x 10-9 M or less;
- comprising the amino acid sequence of SEQ ID NO: 26, a light chain CDR2 comprising koff rate no more than 10-fold higher than the antibody comprising a light chain CDR3 selective mutagenesis position or a hypermutation position, wherein said mutant has a 30, or a mutant thereof having one or more amino acid substitutions at a preferred NO: 28, and a light chain CDR1 comprising the amino acid sequence of SEQ ID NO: SEQ ID NO: 26, a light chain CDR2 comprising the amino acid sequence of SEQ ID comprises a light chain CDR3 comprising the amino acid sequence of

comprising the amino acid sequence of SEQ ID NO: 29; and

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the amino acid sequence of SEQ ID NO: 28, and a light chain CDRI comprising the amino acid sequence of SEQ ID NO: 30.

In yet another embodiment, the invention provides isolated human antibodies, or antigen-binding portions thereof, that neutralize the activity of human IL-12, and at least one additional primate IL-12 selected from the group consisting of baboon IL-12, marmoset IL-12, chimpanzee IL-12, cynomolgus IL-12 and rhesus IL-12, but which do not neutralize the activity of the mouse IL-12.

Recombinant human antibodies of the invention can be isolated by screening of a

Selection of Recombinant Human Antibodies

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The antibody libraries used in this method are preferably scFv libraries prepared 4137; and Barbas et al. (1991) PNAS 88:7978-7982. (1991) Bio/Technology 9:1373-1377; Hoogenboom et al. (1991) Nuc Acid Res 19:4133-(1991) Nature 352:624-628; Gram et al. (1992) PNAS 89:3576-3580; Garrad et al. EMBO J 12:725-734; Hawkins et al. (1992) J Mol Biol 226:889-896; Clackson et al. 246:1275-1281; McCafferty et al., Nature (1990) 348:552-554; Griffiths et al. (1993) Hay et al. (1992) Hum Antibod Hybridomas 3:81-85; Huse et al. (1989) Science PCT Publication No. WO 92/09690; Fuchs et al. (1991) Bio/Technology 9:1370-1372; WO 93/01288; McCafferty et al. PCT Publication No. WO 92/01047; Garrard et al. Winter et al. PCT Publication No. WO 92/20791; Breitling et al. PCT Publication No. libraries can be found in, for example, Kang et al. PCT Publication No. WO 92/18619; and reagents particularly amenable for use in generating and screening antibody display the Stratagene SurfZAPTM phage display kit, catalog no. 240612), examples of methods (e.g., the Pharmacia Recombinant Phuge Antibody System, catalog no. 27-9400-01; and the art. In addition to commercially available kits for generating phage display libraries lymphocytes. Methodologies for preparing and screening such libraries are known in prepared using human VV and VH cDNAs prepared from mRNA derived from human recombinant combinatorial antibody library, preferably a scFv phage display library,

from human VL and VH cDNAs. The scFv antibody libraries are preferably screened using recombinant human IL-12 as the antigen to select human heavy and light chain sequences having a binding activity toward IL-12. To select for antibodies specific for the p35 subunit of IL-12 or the p70 heterodimer, screening assays were performed in the

presence of excess free p40 subunit. Subunit preferences can be determined, for example by, micro-Friguet titration, as described in Example 1.

Once initial human VL and VH segments are selected, "mix and match"

specificity/affinity produced as a result of in vitro affinity maturation. selected. Table 2 (see Appendix A) shows antibodies that displayed altered binding and sequences that exhibit high affinity and a low off rate for IL-12 binding can be mutated VV and VV segments can be reselected and rescreened for binding to hIL-12 mutations have been introduced into the VH and/or VL CDR3 regions. These randomly that the resultant PCR products encode VH and VL segments into which random "spiked" with a random mixture of the four nucleotide bases at certain positions such complimentary to the VH CDR3 or VL CDR3, respectively, which primers have been maturation can be accomplished by amplifying VH and VL regions using PCR primers maturation of antibodies during a natural immune response. This in vitro affinity process analogous to the in vivo somatic mutation process responsible for affinity can be randomly mutated, preferably within the CDR3 region of VH and/or VL, in a constant for hIL-12 binding, the VL and VH segments of the preferred VL/VH pair(s) Example 1). Additionally, to further improve the affinity and/or lower the off rate for IL-12 binding, are performed to select preferred VL/VH pair combinations (see experiments, in which different pairs of the selected VL and VH segments are screened

invention from a recombinant immunoglobulin display library, nucleic acid encoding the selected antibody can be recovered from the phage particle(s) (e.g., from the phage particle(s) (e.g., from the phage genome) and subcloned into other expression vectors by standard recombinant DNA techniques. If desired, the nucleic acid can be further manipulated to create other antibody forms of the invention (e.g., linked to nucleic acid encoding additional immunoglobulin domains, such as additional constant regions). To express a recombinant human antibody isolated by screening of a combinatorial library, the DNA encoding the antibody is cloned into a recombinant expression vector and introduced into a mammalian host cells, as described in further detail in Section IV below.

Methods for selecting human IL-12 binding antibodies by phage display

Following selection, isolation and screening of an anti-hIL-12 antibody of the

technology, and affinity maturation of selected antibodies by random or site-directed mutagenesis of CDR regions are described in further detail in Example 1.

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was similar to the COS-3 germline sequence. COS-3 belongs to the V_H3 family of heavy chain germline sequences selected from the VBASE database, revealed that loe 9 further development. A comparison of the heavy chain variable region of loe 9 with the identified a series of anti-IL-12 antibodies, of which the Joe 9 antibody was selected for As described in Example 1, screening of human VL and VH cDNA libraries

out of approximately 100 VH residues, (i.e., 69-98% amino acid sequence homology between any two germline VH sequences of the $V_{\rm H}3$ family varies from 69-98 residues (1995) Immunology Today, 16, 237-242). The range of amino acid sequence identity high (See e.g., Tomlinson et al. (1992) J. Mol. Biol., 227, 776-798 and Cook et al. antibody sequence, the amino acid sequence identity within the entire $V_H \mathfrak{I}$ family is largest contribution to the germline repertoire. For any given human V_H3 - germline 237-242). The V_H 3 family contains the highest number of members and makes the al. (1992) J. Mol. Biol., 227, 776-798 and Cook et al. (1995) Immunology Today, 16, into seven families, $V_H l - V_H 7$, based on nucleotide sequence homology (Tomlinson et The V_H 3 family is part of the human VH germline repertoire which is grouped germline sequences.

Studies of antibody structures have shown that CDR conformations can be structural features upon the CDRs. the CDR and framework regions of the VH chain. These amino acid residues confer

 $V_{\rm H}$ 3 family members results in certain amino acid residues being present at key sites in sequence homology). The high degree of amino acid sequence homology between the

between any two germline VH sequences). For most pairs of germline sequences there

is at least 80 or more identical amino acid residues, (i.e., at least 80% amino acid

grouped into families of canonical CDR structures based on the key amino acid residues

there is a conservation of amino acid residue identity at the key sites for the CDRI and 196, 901-917 and Chothia et al. (1989) Nature, 342, 877-883). Within the V_H3 family structures with identical key amino acid residues (Chothia et al. (1987) J. Mol. Biol., are similar local CDR conformations in different antibodies that have canonical that occupy certain positions in the CDR and framework regions. Consequently, there

CDR2 canonical structures (Chothia et al. (1992) J. Mol. Biol., 227, 799-817).

The COS-3 germline VH gene, is a member of the V_H3 family and is a variant of the 3-30 (DP-49) germline VH allele. COS-3, differs from Joe9 VH amino acid sequences at only 5 positions. The high degree of amino acid sequence homology between Joe9 VH and COS-3, and between Joe9 VH and the other V_H3 family members also confers a high degree of CDR atructural homology (Chothia et al. (1992) J. Mol. Biol., 227, 799-817; Chothia et al. (1987) J. Mol. Biol., 196, 901-917 and Chothia et al. (1989) Nature, 342, 877-883).

and canonical structural similarity to Joe 9, other V_H3 family members could also be used to generate antibodies that bind to human IL-12. This can be performed, for example, by selecting an appropriate VL by chain-shuffling techniques (Winter et al. (1994) Annual Rev. Immunol., 12, 433-55), or by the graffing of CDRs from a rodent or other human antibody including CDRs from antibodies of this invention onto a V_H3 family framework.

The skilled artisan will appreciate that based on the high amino acid sequence

Incloude sequence homology (Williams et al. (1996) J. Mol. Biol., 264, 220-232). A comparison of the light chain variable region of loe 9 with the light chain germline sequences selected from the VBASE database, revealed that loe 9 was similar to the DPL8 lambda germline. The loe9 VL differs from DPL8 sequence at only four framework positions, and is highly homologous to the framework sequences of the other V\(\frac{1}{2}\) family members. Based on the high amino acid sequence homology and canonical structural similarity to loe 9, other V\(\frac{1}{2}\) family members may also be used to generate arructural similarity to loe 9, other V\(\frac{1}{2}\) family members do be used to generate an appropriate VH by chain-shuffling techniques (Winter et al. Supra, or by the grafting of CDRs from a rodent or other human antibody including CDRs from antibodies of this invention onto a V\(\frac{1}{2}\) family framework.

The methods of the invention are intended to include recombinant antibodies that bind to hIL-12, comprising a heavy chain variable region derived from a member of the V_H3 family of germline sequences, and a light chain variable region derived from a appreciate that any member of the V_H3 family heavy chain sequence can be combined appreciate that any member of the V_H3 family light chain sequence. With any member of the V_A1 family light chain sequence.

Those skilled in the art will also appreciate that DNA sequence polymorphisms that lead to changes in the amino acid sequences of the germline may exist within a population (e.g., the human population). Such genetic polymorphism in the germline sequences may exist among individuals within a population due to natural allelic variations can typically result in 1-5 % variance in the nucleotide sequence of the a gene. Any and all such nucleotide variations and resulting amino acid polymorphisms in germline sequences that are the result of natural allelic variation are intended to be within the scope of the invention.

Accordingly, in one aspect, the invention features an isolated human antibody, or Accordingly, in one aspect, the invention features an isolated human antibody, or

o an antigen-binding portion thereof, which has the following characteristics:

a) that binds to human IL-12 and dissociates from human IL-12 with a

k_{off} rate constant of 0.1 s⁻¹ or less, as determined by surface plasmon resonance, or which

 k_{off} rate constant of 0.1 s⁻¹or less, as determined by surface plasmon resonance, or which inhibits phytohemagglutinin blast proliferation in an *in vit*ro phytohemagglutinin blast proliferation as say (PHA assay) with an IC₅₀ of 1 x 10⁻⁶M or less.

b) has a heavy chain variable region comprising an amino acid sequence selected from a member of the V_H 3 germline family, wherein the heavy chain variable region has a mutation at a contact or hypermutation position with an activity enhancing amino acid residue.

c) has a light chain variable region comprising an amino acid sequence selected from a member of the V_{λ} 1 germline family, wherein the light chain variable region has a mutation at a preferred selective mutagenesis position, contact or hypermutation position with an activity enhancing amino acid residue.

mutation in the heavy chain CDR3.

In a preferred embodiment, the isolated human antibody, or antigen binding has

binding has mutation in the light chain CDR3.

In another preferred embodiment, the isolated human antibody, or antigen

binding has mutation in the heavy chain CDR2.

In another preferred embodiment, the isolated human antibody, or antigen

30 binding has mutation in the light chain CDR2.

In another preferred embodiment, the isolated human antibody, or antigen

binding has mutation in the heavy chain CDR1.

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In another preferred embodiment, the isolated human antibody, or antigen

binding has mutation in the light chain CDRI.

An ordinarily skilled artisan will appreciate that based on the high amino acid

An ordinarily skilled artisan will appreciate that based on the high amino acid sequence similarity between members of the $V_{\rm H}3$ germline family, or between members of the light chain $V_{\lambda}1$ germline family, that mutations to the germlines sequences can provide additional antibodies that bind to human IL-12. Table I (see Appendix A) shows the germline sequences of the $V_{\rm H}3$ family members and demonstrates the

provide additional antibodies that bind to human IL-12. Table I (see Appendix A) shows the germline sequences of the $V_{\rm H}3$ family members and demonstrates the significant sequence homology within the family members. Also shown in Table I are the germline sequences for $V_{\lambda}1$ family members. The heavy and light chain sequences of $V_{\rm H}3$ or of loe 9 are provided as a comparison. Mutations to the germline sequences of $V_{\rm H}3$ or $V_{\lambda}1$ family members may be made, for example, at the same amino acid positions as

those made in the antibodies of the invention (e.g. mutations in Joe 9). The modifications can be performed using standard molecular biology techniques, such as by PCR mutagenesis, targeting individual amino acid residues in the germline sequences, followed by kinetic and functional analysis of the modified antibodies as described herein (e.g., neutralization assays described in Example 3, and by BIAcore analysis, as

described in Example 5). Accordingly, in one aspect, the invention features isolated human antibody, or an

antigen-binding portion thereof, which has the following characteristics:

a) has a heavy chain variable region comprising an amino acid sequence selected from the group consisting of SEQ ID NOs: 595-667, wherein the heavy chain variable region has a mutation at a preferred selective mutagenesis position, contact or hypermutation position with an activity enhancing amino acid residue.

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b) has a light chain variable region comprising an amino acid sequence selected from the group consisting of SEQ ID NOs: 669-675, wherein the light chain variable region has a mutation at a preferred selective mutagenesis position, contact or hypermutation position with an activity enhancing amino acid residue.

An ordinarily skilled artisan will appreciate that based on the high amino acid

sequence similarity between Joe 9 and COS-3 heavy chain germline sequence, and between Joe 9 and DPL8 lambda germline sequence, that other mutations to the CDR regions of these germlines sequences can provide additional antibodies that bind to

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human IL-12. Such methods of modification can be performed using standard molecular

biology techniques as described above.

Accordingly, in one aspect, the invention features isolated human antibody, or an antigen-binding portion thereof, which has the following characteristics:

a) that binds to human IL-12 and dissociates from human IL-12 with a $k_{\rm off}$ rate constant of 0.1s⁻¹or less, as determined by surface plasmon resonance, or which inhibits phytohemagglutinin blast proliferation in an *in vitro* phytohemagglutinin blast

inhibits phytohemagglutinin blast proliferation in an *in vitro* phytohemagglutinin blast proliferation as Novemagglutinin blast proliferation as Novemagglutinin blast proliferation as November 10.50 of 1 x 10.50

b) has a heavy chain variable region comprising the COS-3 germline

amino acid sequence, wherein the heavy chain variable region has a mutation at a preferred selective mutagenesis position, contact or hypermutation position with an activity enhancing amino acid residue.

c) has a light chain variable region comprising the DPL8 germline amino acid sequence, wherein the light chain variable region has a mutation at a preferred selective mutatenesis position contact or hypermutation position with an activity

selective mutagenesis position, contact or hypermutation position with an activity enhancing amino acid residue.

Due to certain amino acid residues occupying key sites in the CDR and framework regions in the light and heavy chain variable region, structural features are conferred at these regions. In particular, the CDR2 and CDR1 regions are subject to homology between family members, these canonical features are present between family members, those canonical features are present between family residues that modifications at the amino acid residues that confer these canonical structures would produce additional antibodies that

bind to IL-12. The modifications can be performed using standard molecular biology

25 techniques as described above.

Accordingly, in another aspect, the invention features an isolated human

antibody, or an antigen-binding portion thereof, which has the following characteristics:

a) that binds to human IL-12 and dissociates from human IL-12 with a

 k_{off} rate constant of 0.1 s⁻¹or less, as determined by surface plasmon resonance, or which in inhibits phytohemagglutinin blast proliferation in an *in vitro* phytohemagglutinin blast proliferation assay (PHA assay) with an IC₅₀ of 1 x 10⁻⁶M or less.

b) has a heavy chain variable region comprising an amino acid sequence selected from a member of the $V_{\rm H}3$ germline family, wherein the heavy chain variable family members, and a CDR1 that is structurally similar to CDR1s from other $V_{\rm H}3$ germline family members, and wherein the heavy chain variable region has a mutation at a preferred selective mutagenesis position, contact or hypermutation position with an activity enhancing amino acid residue;

selected from a member of the $V_{\lambda}I$ germline family, wherein the light chain variable region comprises a CDR2 that is structurally similar to CDR2s from other $V_{\lambda}I$ germline family members, and a CDR1 that is structurally similar to CDR1s from other $V_{\lambda}I$ germline family members, and wherein the light chain variable region has a mutation at a preferred selective mutagenesis position, contact or hypermutation position with an activity enhancing amino acid residue.

c) has a light chain variable region comprising an amino acid sequence

Recombinant human antibodies of the invention have variable and constant

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immunoglobulin framework sequence that differ from the closest germline sequences. chain framework amino acid sequences to identify amino acid residues in the mutated VBASE database) are first compared to the mutated immunoglobulin heavy and light sequences of heavy and light chain encoded by the germline (as found as example in invention can optionally include a backmutation step. To do this, the amino acid "backmutation" of framework residues to the germline configuration). Thus, the present change these amino acid differences back to the true germline sequences (i.e., amplified sequence as compared to the true germline sequence), it may be desirable to framework regions from the true germline configuration (i.e., differences in the sequences obtained by PCR amplification encode amino acid differences in the that occurs during B-cell development. It should be noted that if the "germline" their corresponding germline sequences due to the normal process of somatic mutation which were derived from human donors will contain antibody sequences that differ from 50 human germline immunoglobulin sequences. Also, libraries of recombinant antibodies random mutagenesis or PCR mutagenesis) result in amino acids that are not encoded by from the VBASE database. Mutations to the recombinant human antibodies (e.g., by regions which are homologous to human germline immunoglobulin sequences selected

Then, the appropriate nucleotides of the mutated immunoglobulin sequence are mutated back to correspond to the germline sequence, using the genetic code to determine which nucleotide changes should be made. Mutagenesis of the mutated immunoglobulin framework sequence is carried out by standard methods, such as PCR-mediated such that the PCR product contains the mutations) or site-directed mutagenesis. The role of each amino acid identified as candidate for backmutation should be investigated for a direct or indirect role in antigen binding and any amino acid found after mutation to affect any desirable characteristic of the human antibody should not be included in the affect any desirable characteristic of the human antibody should not be included in the selective mutagenesis approach will not be subject to backmutation. Assays to determine selective mutagenesis approach will not be subject to backmutation. Assays to determine competitive ELISA, in vitro and in vivo neutralization assays and/or (see e.g. Example competitive ELISA, in vitro and in vivo neutralization assays and/or (see e.g. Example of immunohistochemistry with tissue sections from various sources (including human,

To minimize the number of amino acids subject to backmutation those amino acid positions found to be different from the closest germline sequence but identical to the corresponding amino acid in a second germline sequence can remain, provided that the second germline sequence is identical and colinear to the sequence of the human amino acid in question. This would assure that any peptide epitope presented to the immune system by professional antigen presenting cells in a subject treated with the human antibody of the invention would not be foreign but identical to a self-antigen, i.e. occur at any stage of antibody optimization; preferably, backmutation may before or after the selective mutagenesis approach. More preferably, backmutation occurs directly before the selective mutagenesis approach.

primate and/or other species).

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III. Modifications to Preferred Selective Mutagenesis Positions, Contact and/or

30 Hypermutation Positions

Typically, selection of antibodies with improved affinities can be carried out using phage display methods, as described in section II above. This can be

activity of any antibody. system. Moreover, the selective mutagenesis approach can be used to improve the system, it should be noted that this method can also be used with the phage display Mutagenesis Approach was developed to overcome limitations using the phage display to overcome this limitation and is provided by the invention. Although this Selective which does not require phage display affinity maturation of antibodies, was established specificity/affinity. Accordingly, a method termed Selective Mutagenesis Approach methods are limiting in their ability to select antibodies with a highly improved binding phage display methods. Thus, for at least certain antibodies or antigens, phage display Y61). Accordingly, antibodies with even higher affinities could not be selected by IL-12 antibodies, upon attaining a certain level of affinity achieved (i.e., that of antibody particle) when phage display methods were used to improve the affinity of selected anti-(presumably due to additional non-specific interactions between the antigen and phage conditions that would allow equilibrium to be established could not be determined time, preferential binding of higher affinity antibodies to the antigen. Selection methods to work, the antibody-antigen reaction must tend to equilibrium to allow, over libraries containing antibodies of different sequences. However, for these selection accomplished by randomly mutating combinations of CDR residues and generating large

To improve the activity (e.g., affinity or neutralizing activity) of an antibody, ideally one would like to mutate every CDR position in both the heavy and light chains to every other possible amino acid residue. However, since there are, on average, 70 CDR positions within an antibody, such an approach would be very time consuming and labor intensive. Accordingly, the method of the invention allows one to improve the activity of the antibody by mutating only certain selected residues within the heavy and/or light chain CDRs. Furthermore, the method of the invention allows improvement in activity of the antibody without affecting other desirable properties of the antibody in activity of the antibody without affecting other desirable properties of the antibody.

contact with an antigen cannot be accurately predicted based on primary sequence or their positions within the variable region. Nevertheless, alignments of sequences from antibodies with different specificities conducted by Kabat et al. have identified the CDRs as local regions within the variable regions which differ significantly among antibodies (Kabat et al. (1971) Ann. NY Acad. Sci. 190:382-393, , Kabat, E.A., et al.

of Health and Human Services, NIH Publication No. 91-3242). Structural studies have shown that the antigen binding surface is formed by amino acid residues present in the roles or be directly involved in antigen binding. Therefore, for each antigen-antibody pair, amino acid residues within and outside of the CDRs may be important.

The sequence alignment studies by Tomlison et al identified a number of

many somatic mutations may be permitted that do not contribute to antigen binding recovered by restoring only three of the nineteen somatic mutations, demonstrating that hundred fold loss in activity. The full affinity of the anti-Ars antibody could be residues, generating a germline version of the anti-Ars antibody which had a two-(Ars) antibody were simultaneously replaced with their corresponding germline 87:4814-7). Mineteen somatic mutations in a high-affinity anti-p-azophenylarsonate studies on the role of somatic mutations to antibody affinity (Sharon, (1990), PNAS, which do not contact the antigen. This conclusion is further supported by mutational conserved amino acid residues that contact the antigen, and diverse amino acid residues necessarily predict a role of a specific amino acid in antigen binding, and suggest antigen. Furthermore, Tomlison et al. propose that somatic diversity alone does not of an antibody binding site, and potentially provide important interactions with an CDR3 regions, and sections of the light chain CDR3 which are known to lie in the center sites for somatic mutation. However, this analysis excludes the important heavy chain H58, L30, L31, L31A, L50, L53, L91, L92, L93 and L94 were identified as frequent Mol. Biol. 256: 813-817). In particular, positions H31, H31B, H33, H33B, H52B, H56, chain CDR3 which are frequent sites of somatic mutation. (Tomlison et al (1996) J. positions in the heavy and light chain CDR1 and CDR2, and in a portion of the kappa

The result can be explained in part by the nature of antibody diversity itself.

Immature B-cells may produce initially low affinity antibodies that recognize a number of self or non-self antigens. Moreover, antibodies may undergo in the course of affinity maturation sequence variations that may cause self-reactivity. Hypermutation of such low affinity antibodies may serve to abolish self-reactivity ("negative selection") and increase affinity for the foreign antigen. Therefore, the analysis of primary and

activity.

structural data of a large number of antibodies does not provide a method of predicting either (1) the role of somatic hyper-mutation sites in the affinity maturation process versus the process of decreasing affinity towards unwanted antigens, or (2) how a given amino acid contributes to the properties of a specific antigen-antibody pair.

Other attempts to address the role of specific amino acid residues in antigen

recognition were made by analyzing a number of crystal structures of antigen-antibody complexes (MacCallum et al. (1996) J. Mol. Biol. 262: 732-745). The potential role of positions located within and outside the CDRs was indicated. Positions in CDRs involved in antigen binding in more than 10 of 26 analyzed structures included H31, H33, H50, H52, H53, H54, H56, H58, H95, H96, H97, H98 and H100 in the heavy chain and L30A, L32, L91, L92, L93, L94, L96 in the light chain. However, the authors noted that prediction of antigen contacts using these and other structural data may over and under predict contact positions, leading to the speculation that a different strategy may have to be applied to different antigens.

Pini et al. describe randomizing multiple residues in antibody CDR sequences in a large phage display library to rapidly increase antibody affinity (Pini et al. (1998) J. Biol Chem. 273: 21769-21776). However, the high affinity antibodies discussed by Pini et al. had mutations in a total of eight positions, and a reductionary analysis of which changes are absolutely required to improve affinity of the antibody becomes impractical because of the large number of possible combinations to be tested for the smallest

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number of amino acids required.

Furthermore, randomizing multiple residues may not necessarily preserve other desired properties of the antibody. Desirable properties or characteristics of an antibody

are art-recognized and include for example, preservation of non-cross reactivity, e.g.,

with other proteins or human tissues and preservation of antibody sequences that are close to human germline immunoglobulin sequences improvement of neutralization potency. Other desirable properties or characteristics include ability to preserve species expression levels of protein in mammalian cells. The desirable properties or characteristics can be observed or measured using art-recognized techniques including but not limited to ELISA, competitive ELISA, in vitro and in vivo neutralization assays (see e.g. Example 3), immunohistochemistry with tissue sections from different sources

including human, primate or other sources as the need may be, and studies to expression in mammalian cells using transient expression or stable expression.

In addition, the method of Pini et al may introduce more changes than the

minimal number actually required to improve affinity and may lead to the antibodies triggering anti-human-antibody (HAMA) formation in human subjects.

Further, as discussed elsewhere, the phage display as demonstrated here, or other related method including ribosome display may not work appropriately upon reaching certain affinities between antibody and antigen and the conditions required to reach equilibrium may not be established in a reasonable time frame because of additional interactions

The ordinarily skilled artisan may glean interesting scientific information on the origin of antibody diversity from the teachings of the references discussed above. The present invention, however, provides a method for increasing antibody affinity of a specific antigen-antibody pair while preserving other relevant features or desirable characteristics of the antibody. This is especially important when considering the desirability of imparting a multitude of different characteristics on a specific antibody including antigen binding.

including interactions with other phage or ribosome components and the antigen.

If the starting antibody has desirable properties or characteristics which need to be retained, a selective mutagenesis approach can be the best strategy for preserving these desirable properties while improving the activity of the antibody. For example, in the mutagenesis of Y61, the aim was to increase affinity for hIL-12, and to improve the neutralization potency of the antibody while preserving desired properties. Desired

neutralization potency of the antibody while preserving desired properties. Desired properties of Y61 included (1) preservation of non-cross reactivity with other proteins or human tissues, (2) preservation of fine epitope specificity, i.e. recognizing a p40 epitope preferably in the context of the p70 (p40/p35) heterodimer, thereby preventing binding interference from free soluble p40; and (3) generation of an antibody with heavy and light chain amino acid sequences that were as close as possible to their respective germline immunoglobulin sequences.

mutagenesis approach as a strategy for preserving the desirable properties or characteristics of the antibody while improving the affinity and/or neutralization potency. The term "selective mutagenesis approach" is as defined above and includes a

In one embodiment, the method of the invention provides a selective

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method of individually mutating selected amino acid residues. The amino acid residues to be mutated may first be selected from preferred selective mutagenesis positions, then from contact positions, and then from hypermutation positions. The individual selected position can be mutated to at least two other amino acid residue and the effect of the mutation both on the desired properties of the antibody, and improvement in antibody

activity is determined.

The Selective Mutagenesis approach comprises the steps of:

selecting candidate positions in the order 1) preferred selective muselecting candidate positions in the order 1) preferred selective muselecting candidate positions in the order 1) preferred selective muselecting candidate positions in the order 1) preferred selective muselecting candidate positions in the order 1) preferred selective muselecting candidate provide muselecting candidate c

selecting candidate positions in the order 1) preferred selective mutagenesis positions; 2) contact positions 3) hypermutation positions and light chain variable regions of an antibody (CDR3 preferred over CDR2 preferred over CDR1);

individually mutating candidate preferred selective mutagenesis positions,
hypermutation and/or contact positions in the order of ranking, to all possible other
amino acid residues and analyzing the effect of the individual mutations on the activity
of the antibody in order to determine activity enhancing amino acid residues;

if necessary, making stepwise combinations of the individual activity enhancing

amino acid residues and analyzing the effect of the various combinations on the activity of the antibodies; selecting mutant antibodies with activity enhancing amino acid residues and ranking the mutant antibodies based on the location and identity of the amino acid substitutions with regard to their immunogenic potential. Highest ranking is given to mutant antibodies that comprise an amino acid sequence which nearly identical to a variable region sequence that is described in a germline database, or has an amino acid sequence that is comparable to other human antibodies. Lower ranking is given to a variable region sequences or the sequences of other human antibodies. The lowest either germline sequences or the sequences of other human antibodies. The lowest ranking is given to mutant antibodies with an amino acid substitution that has not been ranking is given to mutant antibodies with an amino acid substitution that has not been

ranking is given to mutant antibodies with an amino acid substitution that has not been encountered in a germline sequence or the sequence of another human antibody. As set forth above, mutant antibodies comprising at least one activity enhancing amino acid residue located in CDR3 is preferred over CDR2 which is preferred over CDR1. The CDRs of the heavy chain variable regions are preferred over those of the light chain

variable region.

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The mutant antibodies can also be studied for improvement in activity, e.g. when compared to their corresponding parental antibody. The improvement in activity of the approvement in activity can be determined for example, by neutralization assays, or binding specificity/affinity by surface plasmon resonance analysis (see Example 3). Preferably, the improvement in activity can be at least 2-20 fold higher than the parental antibody. The improvement in activity can be at least "x₁" to "x₂" fold higher than the parental antibody wherein "x₁" and "x₂" are integers between and including 2 to 20, including antibody wherein "x₁" and "x₂" are integers between and including 2 to 20, including tanges within the state range, e.g. 2-15, e.g. 5-10.

studied to determine whether at least one other desirable property has been retained after mutation. For example, with anti-hIL-12 antibodies testing for, (1) preservation of noncross reactivity with other proteins or human tissues, (2) preservation of epitope recognition, i.e. recognizing a p40 epitope preferably in the context of the p70 (p40/p35) heterodimer, thereby preventing binding interference from free soluble p40; and (3)

The mutant antibodies with the activity enhancing amino acid residue also can be

close as possible to their respective germline immunoglobulin sequences, and determining which would be least likely to elicit a human immune response based on the number of differences from the germline sequence. The same observations can be made on an antibody having more than one activity enhancing amino acid residues, e.g. at least three activity enhancing amino acid residues, e.g. at

retention of the desirable property or characteristic has occurred.

An example of the use of a "selective mutagenesis approach", in the mutagenesis

of Y61 is described below. The individual mutations H31S \rightarrow E, L50 \rightarrow Y, or L94G \rightarrow Y

clones were tested, the activity of the combined clone $H31S \rightarrow E + L50 \rightarrow Y + L94G \rightarrow Y$ was no better than $L50 \rightarrow Y + L94G \rightarrow Y$ (1695). Therefore, changing the germline amino acid residue Ser to Glu at position 31 of CDR1 was unnecessary for the improved activity of 1695 over Y61. The selective mutagenesis approach therefore, identified the minimal number of changes that contributed to the final activity, thereby reducing the immunogenic potential of the final antibody and preserving other desired properties of immunogenic potential of the final antibody and preserving other desired properties of

each improved neutralization activity of the antibody. However, when combination

the antibody.

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Isolated DNA encoding the VH and VL produced by the selected mutagenesis approach can be converted into full length antibody chain genes, to Fab fragment genes as to a scFV gene, as described in section IV. For expression of VH and VL regions produced by the selected mutagenesis approach, expression vectors encoding the heavy and light chain can be transfected into variety host cells as described in detail in section IV. Preferred host cells include either prokaryotic host cells, for example, £ coli, or cukaryotic host cells, for example, yeast cells, e.g., \$. cerevisue. Most preferred eukaryotic host cells are mammalian host cells, described in detail in section IV.

The selective mutagenesis approach provides a method of producing antibodies eukaryotic host cells are mutagenesis approach provides a method of producing antibodies.

with improved activities without prior affinity maturation of the antibody by other means. The selective mutagenesis approach provides a method of producing antibodies with improved affinities which have been subject to back mutations. The selective mutagenesis approach also provides a method of improving the activity of affinity matured antibodies.

The skilled artisan will recognize that the selective mutagenesis approach can be used in standard antibody manipulation techniques known in the art. Examples include, but are not limited to, CDR grafted antibodies, chimeric antibodies, scFV fragments, Fab fragments of a full length antibodies and human antibodies from other sources, e.g., transgenic mice.

Rapid large scale mutational analysis of antibodies include in vitro transcription and translation using ribosome display technology (see e.g., Hanes et al., (1997) Proc. Natl. Acad. Sci. 94: 4937-4942; Dall Acqua et al., (1998) Curr. Opin. Struc. Biol. 8: 443-450; He et al., (1997) Nucleic Acid Res. 25: 5132-5134), and U.S. Patent Nos. 5,643,768 and 5,658,754 issued to Kawasaki. The selective mutagenesis approach also provides a method of producing antibodies with improved activities that can be selected

In the methods of the invention, antibodies or antigen binding portions thereof are further modified by altering individual positions in the CDRs of the HCVR and/or LCVR. Although these modifications can be made in phage-displayed antibodies, the method is advantageous in that it can be performed with antibodies that are expressed in other types of host systems, such as bacterial, yeast or mammalian cell expression

using ribosomal display techniques.

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systems. The individual positions within the CDRs selected for modification are based on the positions being a contact and/or hypermutation position.

L52, L53, L55, L91, L92, L93, L94 and L96. Preferred hypermutation positions are selected from the group consisting of H30, H31, H31B, H32, H52, H56, H58, L30, L31, L32, L53 and L93. More preferred amino acid residues (referred to as "preferred serions are selective mutagenesis positions") are both contact and hypermutation positions and are

selective mutagenesis positions") are both contact and hypermutation positions and are selected from the group consisting of H30, H31, H31B, H32, H33, H32, H36, H36, H38, L30, L31, L32, L50, L91, L92, L93, L94. Particularly preferred contact positions are selected from the group consisting of L50 and L94.

Particularly, preferred activity enhancing amino acid residues replace amino acid residues located at positions selected from the group consisting of L50 and L94.

In general, the method of the invention involves selecting a particular preferred

selective mutagenesis position, contact and/or hypermutation position within a CDR of

thereof, randomly mutagenizing that individual position (e.g., by genetic means using a mutagenic oligonucleotide to generate a "mini-library" of modified antibodies), or mutating a position to specific desired amino acids, to identify activity enhancing amino acid residues expressing, and purifying the modified antibodies (e.g., in a non-phage display host system), measuring the activity of the modified antibodies for antigen (e.g., by measuring k_{off} rates by BIAcore analysis), repeating these steps for other CDR by measuring as necessary, and combining individual mutations shown to have improved positions, as necessary, and combining individual mutations shown to have improved

H31B, H32, H33, H35, H50, H52, H52A, H53, H54, H56, H58, H95, H96, H97, H98, H101, L30, L31, L32, L34, L50, L52, L53, L53, L91, L92, L93, L94 and L96. Preferred hypermutation positions are selected from the group consisting of H30, H31, H31B,

ordinarily skilled artisan using art-recognized techniques.

Preferred contact positions are selected from the group consisting of H30, H31, H31B, H32, H33, H35, H52, H52A, H53, H54, H56, H58, H95, H97, H98,

to the parent antibody, or antigen-binding portion thereof, is obtained. Preferably, the selected antibody or antibodies have an improved activity without loss or with retention of at least one desirable characteristic or property of the parental antibody as described above. The desirable characteristic or property can be measured or observed by the ordinarily skilled artisan using art-recognized techniques.

- portions thereof, relative to the parent antibody or antigen-binding portion thereof, until an antibody, or antigen-binding portion thereof, with an improved activity, relative to the parent antibody, or antigen-binding portion thereof, is obtained. Preferably, the selected antibody or antibodies have an improved activity without loss or with retention as described antibody or antibodies have an improved activity without loss or with retention
- 20 or antigen-binding portions thereof; and goralises, or antigen-binding goralinging the activity of the combination antibodies, or antigen-binding
- individual mutations shown to have improved activity, to form combination antibodies,
- f) combining, in the parent antibody. or antigen-binding portion thereof,
 - selective mutagenesis position, contact or hypermutation position;
 - portions thereof, relative to the parent antibody or antigen-binding portion thereof;
 e) optionally, repeating steps a) through d) for at least one other preferred
- contact or hypermutation position to at least two other amino acid residues to thereby create a panel of mutated antibodies, or antigen-binding portions thereof;

 d) evaluating the activity of the panel of mutated antibodies, or antigen-binding
- position, contact or hypermutation position;c) individually mutating said selected preferred selective mutagenesis position,
 - position, or 3) hypermutation position within a complementarity determining region (CDR) for mutation, thereby identifying a selected preferred selective mutagenesis
 - b) selecting in order a 1) preferred selective mutagenesis position, 2) contact
 - a) providing a parent antibody or antigen-binding portion thereof;
- binding portion thereof.

 Accordingly, in one embodiment, the invention provides a method for improving the activity of an antibody, or antigen-binding portion thereof, comprising:
 - activity and testing whether the combination(s) generate an antibody with even greater activity (e.g., affinity or neutralizing potency) than the parent antibody, or antigen-

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ς In another embodiment, the invention provides a method for improving the preferred contact positions are selected from the group consisting of L50 and L94. H33, H52, H56, H58, L30, L31, L32, L50, L91, L92, L93 and L94. Particularly mutagenesis positions are selected from the group consisting of H30, H31, H31B, H32, H32, H52, H56, H58, L30, L31, L32, L53 and L93. More preferred selective

a) providing a parent antibody or antigen-binding portion thereof; activity of an antibody, or antigen-binding portion thereof, comprising: 10

- b) selecting a preferred selective mutagenesis position, contact or hypermutation
- position within a complementarity determining region (CDR) for mutation;
- create a panel of mutated antibodies, or antigen-binding portions thereof; contact or hypermutation position to at least two other amino acid residues to thereby c) individually mutating said selected preferred selective mutagenesis position,
- thereby identifying an activity enhancing amino acid residue; portions thereof, relative to the parent antibody or antigen-binding portion thereof, d) evaluating the activity of the panel of mutated antibodies, or antigen-binding
- selective mutagenesis position, contact or hypermutation position; e) optionally, repeating steps a) through d) for at least one other preferred
- form combination antibodies, or antigen-binding portions thereof; and individual activity enhancing amino acid residues shown to have improved activity, to f) combining, in the parent antibody, or antigen-binding portion thereof. two
- until an antibody, or antigen-binding portion thereof, with an improved activity, relative antibody or antigen-binding portion thereof; portions thereof with two activity enhancing amino acid residues, relative to the parent

g) evaluating the activity of the combination antibodies, or antigen-binding

to the parent antibody, or antigen-binding portion thereof, is obtained.

mutagenesis positions are selected from the group consisting of H30, H31, H31B, H32, H32, H52, H56, H58, L30, L31, L32, L53 and L93. More preferred preferred selective hypermutation positions are selected from the group consisting of H30, H31, H31B, H101, L30, L31, L32, L34, L50, L52, L53, L55, L91, L92, L93, L94 and L96. Preferred H31B' H35' H33' H32' H20' H25' H25Y' H23' H24' H26' H28' H62' H62' H64' Preferred contact positions are selected from the group consisting of H30. H31,

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In another embodiment, the invention provides a method for improving the preferred contact positions are selected from the group consisting of L50 and L94. H33, H52, H56, H58, L30, L31, L32, L50, L91, L92, L93 and L94. Particularly

activity of an antibody, or antigen-binding portion thereof, comprising:

b) selecting a preferred selective mutagenesis position, contact or hypermutation a) providing a parent antibody or antigen-binding portion thereof;

position within a complementarity determining region (CDR) for mutation;

c) individually mutating said selected preferred selective mutagenesis position,

contact or hypermutation position to at least two other amino acid residues to thereby

d) evaluating the activity of the panel of mutated antibodies, or antigen-binding create a panel of mutated antibodies, or antigen-binding portions thereof;

portions thereof, relative to the parent antibody or antigen-binding portion thereof,

thereby identifying an activity enhancing amino acid residue;

e) optionally, repeating steps a) through d) for at least one other preferred

selective mute genesis position, contact or hypermutation position;

f) combining, in the parent antibody, or antigen-binding portion thereof, three

individual activity enhancing amino acid residues shown to have improved activity, to

form combination antibodies, or antigen-binding portions thereof; and

g) evaluating the activity of the combination antibodies, or antigen-binding

portions thereof with two activity enhancing amino acid residues, relative to the parent

antibody or antigen-binding portion thereof;

to the parent antibody, or antigen-binding portion thereof, is obtained. until an antibody, or antigen-binding portion thereof, with an improved activity, relative

Preferably, the activity enhancing amino acid residue replaces amino acid

residues located at positions selected from the group consisting of H30, H31, H31B,

H35' H32' H32' H20' H25' H25∀' H23' H2¢' H2¢' H3¢' H36' H36' H36' H101'

F30' F31' F35' F34' F20' F25' F23' F22' F61' F65' F63' F64 and F66.

Following mutagenesis of individual selected positions, mutated clones can be

sequenced to identify which amino acid residues have been introduced into the selected

position in each clone. A small number of clones (e.g., about 24) can be selected for

sequencing, which statistically should yield 10-15 unique antibodies, whereas larger

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numbers of clones (e.g., greater than 60) can be sequenced to ensure that antibodies with every possible substitution at the selected position are identified.

In one embodiment, contact and/or hypermutation positions within the CDR3

regions of the heavy and/or light chains are first selected for mutagenesis. However, for antibodies that have already been affinity matured in vitro by random mutagenesis of the CDR3 regions via phage display selection, it may be preferably to first select contact and/or hypermutation positions within CDR1 or CDR2 of the heavy and/or light chain. In a more preferred embodiment, preferred selective mutagenesis positions

within the CDR3 regions of the heavy and/or light chains are first selected for mutagenesis. However, for antibodies that have already been affinity matured in vitro by random mutagenesis of the CDR3 regions via phage display selection, it may be preferably to first select preferred selective mutagenesis positions within CDR1 or CDR2 of the heavy and/or light chain.

In another preferred embodiment, the optimization of a selected antibody by the

retained characteristic or property discussed elsewhere). for increased affinity, neutralization potency (and possibly also for at least one other each (preferably 5-14 other amino acids) and the resulting antibodies are characterized H54, H95, H96, H97, H98, L30A and L96 are mutated to at least 2 other amino acids will be selected for selective mutagenesis from the group consisting of H35, H50, H53, activity (including affinity and/or neutralization potency), additional amino acid residues mutagenesis positions does not result in an combination antibody which meets the target multiple activity enhancing amino acids replacing amino acids in preferred selective affinity or neutralization potency at all or sufficiently and if even the combination of mutation of a single preferred selective mutagenesis position does not increase the also for at least one other retained characteristic or property discussed elsewhere). If a antibodies are characterized for increased affinity, neutralization potency (and possibly least 2 other amino acids each (preferably 5-14 other amino acids) and the resulting H33, H52, H56, H58, L30, L31, L32, L50, L91, L92, L93, L94 are mutated first to at mutagenesis positions selected from the group consisting of H30, H31, H31B, H32, selective mutagenesis approach is done sequentially as follows: preferred selective

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acids) and the resulting antibodies are characterized for increased affinity, neutralization L31A and are mutated to at least 2 other amino acids each (preferably 5-14 other amino will be selected for selective mutagenesis from the group consisting of H33B, H52B, (including affinity and/or target neutralization potency), additional amino acid residues positions does not result in an combination antibody which meets the targeted activity combination of multiple activity enhancing amino acids replacing amino acids in those (including affinity and/or neutralization potency) at all or not sufficiently and if even the H35, H50, H53, H54, H95, H96, H97, H98, L30A and L96 does not increase the activity If a mutation of a single amino acid residue selected from the group consisting of

It should be understood that the sequential selective mutagenesis approach may discussed elsewhere).

potency (and possibly also for at least one other retained characteristic or property

other desired characteristics and are therefore not acceptable, the remaining CDR neutralization potency) and/or if the identified activity enhancing amino acids also affect combination antibody still do not meet the targets set for activity (including affinity and preselected positions has identified activity enhancing amino acids residues but the (including affinity and neutralization potency) has been identified. If mutagenesis of the end at any of the steps outline above as soon as an antibody with the desired activity

residues may be subjected to mutagenesis (see section IV).

The method of the invention can be used to improve activity of an antibody, or

predetermined affinity and/or neutralization potency, and/or a desired property or antigen binding portion thereof, to reach a predetermined target activity (e.g. a

antibody, or antigen-binding portion thereof, to attain a predetermined target activity, Accordingly, the invention provides a method of improving the activity of an characteristic).

consisting of H30, H31, H31B, H32, H32, H52, H56, H58, L30, L31, L32, L50, L91, b) selecting a preferred selective mutagenesis position selected from group a) providing a parent antibody a antigen-binding portion thereof;

L92, L93, L94.

comprising:

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c) individually mutating the selected preferred selective mutagenesis position to at least two other amino acid residues to hereby create a first panel of mutated antibodies, or antigen binding portions thereof;

d) evaluating the activity of the first panel of mutated antibodies, or antigen binding portions thereof to determined if mutation of a single selective mutagenesis position produces an antibody or antigen binding portion thereof v

mutagenesis position produces an antibody or antigen binding portion thereof with the predetermined target activity or a partial target activity;

e) combining in a stepwise fashion, in the parent antibody, or antigen binding

e) combination antibodies, or antigen binding portions thereof.

f) evaluating the activity of the combination antibodies, or antigen binding portions thereof to determined if the combination antibodies, or antigen binding portions thereof have the predetermined target activity or a partial target activity.

g) if steps d) or f) do not result in an antibody or antigen binding portion thereof

having the predetermined target activity, or result an antibody with only a partial activity, additional amino acid residues selected from the group consisting of H35, H50, H54, H95, H96, H97, H98, L30A and L96 are mutated to at least two other amino acid residues to thereby create a second panel of mutated antibodies or antigen-binding notitions thereof:

portions thereof;

20 h) evaluating the activity of the second panel of mutated antibodies or antigen

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binding portions thereof, to determined if mutation of a single amino acid residue selected from the group consisting of H35, H50, H53, H54, H95, H96, H97, H98, L30A and L96 results an antibody or antigen binding portion thereof, having the predetermined target activity or a partial activity;

25 i) combining in stepwise fashion in the parent antibody, or antigen-binding portion thereof, individual mutations of step g) shown to have an improved activity, to

form combination antibodies, or antigen binding portions thereof;

j) evaluating the activity of the combination antibodies or antigen binding

portions thereof, to determined if the combination antibodies, or antigen binding portions thereof have the predetermined target activity or a partial target activity;

k) if steps h) or j) do not result in an antibody or antigen binding portion thereof having the predetermined target activity, or result in an antibody with only a partial activity, additional amino acid residues selected from the group consisting of H33B, H52B and L31A are mutated to at least two other amino acid residues to thereby create a third panel of mutated antibodies or antigen binding portions thereof;

l) evaluating the activity of the third panel of mutated antibodies or antigen binding portions thereof, to determine if a mutation of a single amino acid residue selected from the group consisting of H33B, H52B and L31A resulted in an antibody or antigen binding portion thereof, having the predetermined target activity or a partial activity;

m) combining in a stepwise fashion in the parent antibody, or antigen binding portion thereof, individual mutation of step k) shown to have an improved activity, to form combination antibodies, or antigen binding portions, thereof;

n) evaluating the activity of the combination antibodies or antigen-binding

portions thereof, to determine if the combination antibodies, or antigen binding portions thereof have the predetermined target activity to thereby produce an antibody or antigen binding portion thereof with a predetermined target activity.

A number of mutagenesis methods can be used, including PCR assembly, Kunkel (dut-ung-) and thiophosphate (Amersham Sculptor kit) oligonucleotide-directed

mutagenesis.

A wide variety of host expression systems can be used to express the mutated antibodies, including bacterial, yeast, baculoviral and mammalian expression systems (as well as phage display expression systems). An example of a suitable bacterial expression vector is pUC119(Sfi). Other antibody expression systems are known in the art and/or are described below in section IV.

The modified antibodies, or antigen binding portions thereof, produced by the method of the invention can be identified without the reliance on phage display methods for selection. Accordingly, the method of the invention is particularly advantageous for improving the activity of a recombinant parent antibody or antigen-binding portion thereof, that was obtained by selection in a phage-display system but whose activity cannot be further improved by mutagenesis in the phage-display system.

Accordingly, in another embodiment, the invention provides a method for improving the affinity of an antibody, or antigen-binding portion thereof, comprising:

a) providing a recombinant parent antibody or antigen-binding portion thereof; that was obtained by selection in a phage-display system but whose activity cannot be

- further improved by mutagenesis in said phage-display system;

 b) selecting a preferred selective mutagenesis position, contact or hypermutation position within a complementarity determining region (CDR) for mutation, thereby
- position within a complementarity determining region (CDR) for mutation, thereby identifying a selected contact or hypermutation position;
- contact or hypermutation position to at least two other amino acid residues to thereby
- contact or hypermutation position to at least two other amino acid residues to thereby create a panel of mutated antibodies. or antigen-binding portions thereof, and expressing said panel in a non-phage display system;
- said panel in a non-phage display system;

 d) evaluating the activity of the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof;
- e) optionally repeating steps b) through d) for at least one other preferred selective mutagenesis position, contact or hypermutation position;
- f) combining, in the parent antibody, or antigen-binding portion thereof, individual mutations shown to have improved activity, to form combination antibodies, or antigen-binding portions thereof, and

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- g) evaluating the activity of the combination antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof, with an improved activity, relative to the parent antibody, or antigen-binding portion thereof, is obtained.

 Preferred contact positions are selected from the group consisting of H30, H31,
- H101, L30, L31, L32, L34, L50, L52, L53, L55, L91, L92, L93, L94 and L96. Preferred hypermutation positions are selected from the group consisting of H30, H31, H31, H32, H52, H56, H58, L30, L31, L32, L53 and L93. More preferred preferred selective mutagenesis positions are selected from the group consisting of H30, H31, H31B, H32, mutagenesis positions are selected from the group consisting of H30, H31, H31B, H32, H33, H52, H56, H58, L30, L31, L32, L50, L91, L92, L93 and L94. Particularly preferred contact positions are selected from the group consisting of L50 and L94.

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With available methods it is not possible or it is extremely laborious to derive an antibody with increased binding affinity and neutralization potency while retaining other properties or characteristics of the antibodies as discussed above. The method of this invention, however, can readily identify such antibodies. The antibodies subjected to the method of this invention can come from any source.

Therefore, in another embodiment, the invention provides a method for improving the activity of an antibody, or antigen-binding portion thereof, comprising:

a) providing a recombinant parent antibody or antigen-binding portion thereof;

b) selecting a preferred selective mutagenesis position, contact or hypermutation

- position within a complementarity determining region (CDR) for mutation, thereby identifying a selected preferred selective mutagenesis position, contact or hypermutation
- position;

 c) individually mutating said selected preferred selective mutagenesis position, contact or hypermutation position to at least two other amino acid residues to thereby
- contact or hypermutation position to at least two other amino acid residues to thereby create a panel of mutated antibodies, or antigen-binding portions thereof and expressing said panel in an appropriate expression system:
- said panel in an appropriate expression system;

 d) evaluating the activity of the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof,
- thereby identifying an activity enhancing amino acid residue;

 e) evaluating the panel of mutated antibodies, or antigen-binding portions

thereof, relative to the parent antibody or antigen-binding portion thereof for at least one

- other property or characteristics, wherein the property or characteristic is one that needs to be retained in the antibody; until an antibody, or antigen-binding portion thereof, with an improved activity and at
- Linding portion thereof, is obtained.
- In a preferred embodiment, the contact positions are selected from the group consisting of H30, H31, H31B, H32, H33, H35, H50, L52, L53, L53, L53, L54, L50, L52, L53, L53, L54, L50, L54, L50, L52, L53, L55, L91, L92, L93, L94 and L96 and the other characteristic is selected from 1) preservation of non-crossreactivity with other proteins or human tissues. 2) preservation of non-crossreactivity with other proteins or human tissues. 2) preservation of enitone
- crossreactivity with other proteins or human tissues, 2) preservation of epitope recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35

heterodimer preventing binding interference from free, soluble p40 and/or 3) to produce an antibody with a close to germline immunoglobulin sequence.

In another preferred embodiment, the hypermutation positions are selected from the group consisting of H30, H31, H31B, H32, H52, H56, H58, L30, L31, L32, L53 and L93 and the other characteristic is selected from 1) preservation of non-crossreactivity

with other proteins or human tissues, 2) preservation of epitope recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 heterodimer preventing binding interference from free, soluble p40 and/or 3) to produce an antibody with a close to germline immunoglobulin sequence.

In a more preferred embodiment the residues for selective mutagenesis are selected from the preferred selective mutagenesis positions from the group consisting of H30, H31, H31B, H32, H33, H52, H56, H58, L30, L31, L32, L50, L91, L92, L93, L94 and the other characteristic is selected from 1) preservation of non-crossreactivity with other proteins or human tissues, 2) preservation of epitope recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 heterodimer preventing binding interference from free, soluble p40 and/or 3) to produce an antibody with a close to

germline immunoglobulin sequence.

In a more preferred embodiment, the contact positions are selected from the

group consisting of L50 and L94 and the other characteristic is selected from 1) preservation of non-crossreactivity with other proteins or human tissues, 2) preservation of epitope recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 heterodimer preventing binding interference from free, soluble p40 and/or 3) to produce an antibody with a close to germline immunoglobulin sequence.

If therefore, the affinity of an antibody for a specific antigen should be improved,

but where the phage display (or related system including ribosome display) method is no longer applicable, and other desirable properties or characteristics should be retained, the method of the invention can be used. Accordingly, in another embodiment, the invention provides a method for improving the activity of an antibody, or antigen-binding portion thereof, comprising:

a) providing a recombinant parent antibody or antigen-binding portion thereof; that was obtained by selection in a phage-display system; further improved by mutagenesis in said phage-display system;

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b) selecting a preferred selective mutagenesis position, contact or hypermutation position within a complementarity determining region (CDR) for mutation, thereby identifying a selected preferred selective mutagenesis position, contact or hypermutation

- position;

 contact or hypermutation position to at least two other amino acid residues to thereby
 create a panel of mutated antibodies, or antigen-binding portions thereof, and expressing
- create a panel of mutated antibodies, or antigen-binding portions thereof, and expressing said panel in a non-phage display system;
- and paner in a non-phage dispital system,

 d) evaluating the activity of the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof
- thereby identifying an activity enhancing amino acid residue;

 e) evaluating the panel of mutated antibodies, or antigen-binding portions
- thereof, relative to the parent antibody or antigen-binding portion thereof for at least one other property or characteristic, wherein the property or characteristic is one that needs to be retained, until an antibody, or antigen-binding portion thereof, with an improved activity and at least one retained property or characteristic, relative to the parent
- activity and at least one retained property or characteristic, relative to the patent antibody, or antigen-binding portion thereof, is obtained.

 f) optionally, repeating steps a) through e) for at least one other preferred
- selective mutagenesis position, contact or hypermutation position;
 20 g) combining, in the parent antibody, or antigen-binding portion thereof, at least
- two individual activity enhancing amino acid residues shown to have improved activity and at least one retained property or characteristic, to form combination antibodies, or antigen-binding portions thereof; and

h) evaluating the activity of the combination antibodies, or antigen-binding

- portions thereof, relative to the parent antibody or antigen-binding portion thereof, until an antibody, or antigen-binding portion thereof, with an improved activity and at least one retained other property or characteristic, relative to the parent antibody, or

antigen-binding portion thereof, is obtained.

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heterodimer preventing binding interference from free, soluble p40 and/or 3) to produce recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 crossreactivity with other proteins or human tissues, 2) preservation of epitope

with other proteins or human tissues, 2) preservation of epitope recognition, i.e. L93 and the other characteristic is selected from 1) preservation of non-crossreactivity the group consisting of H30, H31, H31B, H32, H52, H56, H58, L30, L31, L32, L53 and In another preferred embodiment, the hypermutation positions are selected from an antibody with a close to germline immunoglobulin sequence.

preventing binding interference from free, soluble p40 and/or 3) to produce an antibody recognizing p40 epitope preferably in the context of the p70 p40/p35 heterodimer

In a more preferred embodiment the residues for selective mutagenesis are with a close to germline immunoglobulin sequence.

interference from free, soluble p40 and/or 3) to produce an antibody with a close to p40 epitope preferably in the context of the p70 p40/p35 heterodimer preventing binding other proteins or human tissues, 2) preservation of epitope recognition, i.e. recognizing and the other characteristic is selected from 1) preservation of non-crossreactivity with H30' H31' H31B' H35' H33' H25' H26' H28' T30' T31' T35' T20' T51' T65' T64 selected from the preferred selective mutagenesis positions from the group consisting of

germline immunoglobulin sequence.

In a more preferred embodiment, the contact positions are selected from the

p40/p35 heterodimer preventing binding interference from free, soluble p40 and/or 3) to of epitope recognition, i.e. recognizing p40 epitope preferably in the context of the p70 preservation of non-crossreactivity with other proteins or human tissues, 2) preservation group consisting of L50 and L94 and the other characteristic is selected from 1)

activity of an antibody, or antigen-binding portion thereof, comprising: In another embodiment, the invention provides a method for improving the

produce an antibody with a close to germline immunoglobulin sequence.

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further improved by mutagenesis in said phage-display system; that was obtained by selection in a phage-display system but whose activity cannot be a) providing a recombinant parent antibody or antigen-binding portion thereof;

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position within a complementarity determining region (CDR) for mutation, thereby b) selecting a preferred selective mutagenesis position, contact or hypermutation

c) individually mutating said selected preferred selective mutagenesis position, identifying a selected contact or hypermutation position;

said panel in a non-phage display system; create a panel of mutated antibodies, or antigen-binding portions thereof, and expressing contact or hypermutation position to at least two other amino acid residues to thereby

d) evaluating the activity of the panel of mutated antibodies. or antigen-binding

thereby identifying an activity enhancing amino acid residue; portions thereof, relative to the parent antibody or antigen-binding portion thereof

to be retained, until an antibody, or antigen-binding portion thereof, with an improved other property or characteristic, wherein the property or characteristic is one that needs thereof, relative to the parent antibody or antigen-binding portion thereof for at least one e) evaluating the panel of mutated antibodies, or antigen-binding portions

S١ activity and at least one retained property or characteristic, relative to the parent

In a preferred embodiment, the contact positions are selected from the group antibody, or antigen-binding portion thereof, is obtained.

L94 and L96 and the other characteristic is selected from 1) preservation of non-H92' H96' H91' H98' H101' F30' F31' F37' F34' F20' F35' F23' F32' F93' F93' consisting of H30, H31, H31B, H32, H33, H35, H50, H52, H52A, H53, H54, H56, H58,

an antibody with a close to germline immunoglobulin sequence. heterodimer preventing binding interference from free, soluble p40 and/or 3) to produce recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 crossreactivity with other proteins or human tissues, 2) preservation of epitope

In another preferred embodiment, the hypermutation positions are selected from

preventing binding interference from free, soluble p40 and/or 3) to produce an antibody recognizing p40 epitope preferably in the context of the p70 p40/p35 heterodimer with other proteins or human tissues, 2) preservation of epitope recognition, i.e. L93 and the other characteristic is selected from 1) preservation of non-crossreactivity the group consisting of H30, H31, H31B, H32, H52, H56, H58, L30, L31, L32, L53 and

with a close to germline immunoglobulin sequence.

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interference from free, soluble p40 and/or 3) to produce an antibody with a close to p40 epitope preferably in the context of the p70 p40/p35 heterodimer preventing binding other proteins or human tissues. 2) preservation of epitope recognition, i.e. recognizing and the other characteristic is selected from 1) preservation of non-crossreactivity with H30, H31, H31B, H32, H33, H52, H56, H58, L30, L31, L32, L50, L91, L92, L94, L94 selected from the preferred selective mutagenesis positions from the group consisting of In a more preferred embodiment the residues for selective mutagenesis are

In a more preferred embodiment, the contact positions are selected from the germline immunoglobulin sequence.

produce an antibody with a close to germline immunoglobulin sequence. p40/p35 heterodimer preventing binding interference from free, soluble p40 and/or 3) to of epitope recognition, i.e. recognizing p40 epitope preferably in the context of the p70 preservation of non-crossreactivity with other proteins or human tissues, 2) preservation group consisting of L50 and L94 and the other characteristic is selected from 1)

activity of an antibody, or antigen-binding portion thereof, comprising: In another embodiment, the invention provides a method for improving the

that was obtained by selection in a phage-display system but whose activity cannot be a) providing a recombinant parent antibody or antigen-binding portion thereof;

b) selecting a preferred selective mutagenesis position, contact or hypermutation further improved by mutagenesis in said phage-display system;

identifying a selected contact or hypermutation position; position within a complementarity determining region (CDR) for mutation, thereby

create a panel of mutated antibodies, or antigen-binding portions thereof, and expressing contact or hypermutation position to at least two other amino acid residues to thereby c) individually mutating said selected preferred selective mutagenesis positions,

d) evaluating the activity of the panel of mutated antibodies, or antigen-binding said panel in a non-phage display system;

thereby identifying an activity enhancing amino acid residue; portions thereof, relative to the parent antibody or antigen-binding portion thereof

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other property or characteristic, wherein the property or characteristic is one that needs thereof, relative to the parent antibody or antigen-binding portion thereof for at least one

e) evaluating the panel of mutated antibodies, or antigen-binding portions

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to be retained, until an antibody, or antigen-binding portion thereof, with an improved activity and at least one retained characteristic, relative to the parent antibody, or antigen-binding portion thereof, is obtained.

- f) optionally, repeating steps a) through e) for at least one other preferred selective mutagenesis position, contact or hypermutation position;
- g) combining, in the parent antibody, or antigen-binding portion thereof, at least two individual activity enhancing amino acid residues shown to have improved activity and at least on retained other characteristic, to form combination antibodies, or antigenbinding portions thereof; and
- h) evaluating the activity of the combination antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof, with an improved activity and at least one retained property or characteristic, relative to the parent antibody, or antigenbinding portion thereof, is obtained.
- In a preferred embodiment, the contact positions are selected from the group consisting of H30, H31, H31B, H32, H35, H36, H50, H52, H52A, H53, H54, H56, H58, H95, H96, H97, H98, H101, L30, L31, L32, L34, L50, L52, L53, L53, L55, L91, L92, L93, L94 and L96 and the other characteristic is selected from 1) preservation of non-recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 heterodimer preventing binding interference from free, soluble p40 and/or 3) to produce
- an antibody with a close to germline immunoglobulin sequence.

 In another preferred embodiment, the hypermutation positions are selected from the group consisting of H30, H31, H31B, H32, H52, H56, H58, L30, L31, L32, L53 and
- L93 and the other characteristic is selected from 1) preservation of non-crossreactivity with other proteins or human tissues, 2) preservation of epitope recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 heterodimer preventing binding interference from free, soluble p40 and/or 3) to produce an antibody with a close to germline immunoglobulin sequence.
- selected from the preferred selective mutagenesis positions from the group consisting of H30, H31, H31B, H32, H33, H52, H56, H58, L30, L31, L32, L50, L91, L92, L93, L94

In a more preferred embodiment the residues for selective mutagenesis are

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and the other characteristic is selected from 1) preservation of non-crossreactivity with other proteins or human tissues, 2) preservation of epitope recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 heterodimer preventing binding interference from free, soluble p40 and/or 3) to produce an antibody with a close to germline immunoglobulin sequence.

In a more preferred embodiment, the contact positions are selected from the

group consisting of L50 and L94 and the other characteristic is selected from 1) preservation of non-crossreactivity with other proteins or human tissues, 2) preservation of epitope recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 heterodimer preventing binding interference from free, soluble p40 and/or 3) to produce an antibody with a close to germline immunoglobulin sequence.

IV. Modifications of other CDR residues

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Ultimately, all CDR residues in a given antibody-antigen pair identified by any means to be required as activity enhancing amino acid residues and/or required directly or indirectly for binding to the antigen and/or for retaining other desirable properties or characteristics of the antibody. Such CDR residues are referred to as "preferred selective mutagenesis residues can be identified also by other means including preferred selective mutagenesis residues can be identified also by other means including co-crystallization of antibody and antigen and molecular modeling.

If the preferred attempts to identify activity enhancing amino acids focussing on It the preferred attempts to identify activity enhancing amino acids focussing on

the preferred selective mutagenesis positions, contact or hypermutation positions described above are exhausted, or if additional improvements are required, the remaining CDR residues may be modified as described below. It should be understood positions according to the embodiments discussed above but may require further improvements. Therefore, in another embodiment, the invention provides a method for improving the activity of an antibody, or antigen-binding portion thereof, comprising:

a) providing a parent antibody or antigen-binding portion thereof,

(CDR) for mutation other than H30, H31, H31B, H32, H33, H35, H50, H52, H52A,

b) selecting an amino acid residue within a complementarity determining region

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H23<mark>' H24' H26' H28' H62' H62' H6</mark>3' H68' H101' T30' T31' T35' T34' T20' T23' T23'

- L55, L91, L92, L93, L94 and L96;
 c) individually mutating said selected position e.g., to at least two other amino acid residues to thereby create a mutated antibody or a panel of mutated antibodies, or
- acid residues to thereby create a mutated antibody or a panel of mutated antibodies, or antigen-binding portions thereof;
- d) evaluating the activity of the mutated antibody or the parent antibody or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereby identifying an activity enhancing amino acid residue;
- e) evaluating the mutated antibody or the panel of mutated antibodies, or antigenbinding portions thereof, relative to the parent antibody or antigen-binding portion thereof, with an improved activity, relative to the parent antibody, or antigen-binding portion thereof, with an improved activity, relative to the parent antibody, or antigen-binding portion thereof, is obtained.
- non-crossreactivity with other proteins or human tissues, 2) preservation of epitope recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 heterodimer preventing binding interference from free, soluble p40 and/or 3) to produce an antibody with a close to germline immunoglobulin sequence.

 If mutagenesis of a single residue is not sufficient other residues can be included:

Preferably, the other characteristic or property is selected from 1) preservation of

- If mutagenesis of a single residue is not sufficient other residues can be included; therefore, in another embodiment, the invention provides a method for improving the activity of an antibody, or antigen-binding portion thereof, comprising:
- a) providing a parent antibody or antigen-binding portion thereof;

 b) selecting an amino acid residue within a complementarity determining region

 complementation other than H30, H31, H31B, H32, H33, H35, H50, H52, H52A, H53, H54, H56, H58, H95, H96, H97, H98, H101, L30, L31, L32, L34, L50, L52, L53,
- c) individually mutating said selected position to at least two other amino acid residues to thereby create a panel of mutated antibodies, or antigen-binding portions

thereof;

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d) evaluating the activity of the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof, thereby identifying an activity enhancing amino acid residue;

- e) repeating steps b) through d) for at least one other CDR position which is neither the position selected under b) nor a position at H30, H31, H31B, H32, H33, H35, H50, H52, H52A, H53, H54, H56, H58, H95, H96, H97, H98, H101, L30, L31, L32, L34, L50, L52, L53, L53, L91, L92, L93, L94 and L96;
- f) combining, in the parent antibody, or antigen-binding portion thereof, at least two individual activity enhancing amino acid residues shown to have improved activity, to form combination antibodies, or antigen-binding portions thereof; and
- g) evaluating the activity of the combination antibodies, or antigen-binding portions thereof with two activity enhancing amino acid residues, relative to the parent antibody or antigen-binding portion thereof until an antibody, or antigen-binding portion thereof, with an improved activity, relative to the parent antibody, or antigen-binding portion portion thereof, is obtained.

If the preferred attempts to identify activity enhancing amino acids focussing on the contact or hypermutation positions described above are exhausted, or if additional improvements are required, and the antibody in question can not further be optimized by mutagenesis and phage display (or related ribosome display) methods the remaining antibody could already be modified as described below. It should be understood that the antibody could already be modified in any one or more preferred selective mutagenesis position, contact or hypermutation positions according to the embodiments discussed above but may require further improvements.

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Therefore, in another embodiment, the invention provides a method for improving the activity of an antibody, or antigen-binding portion thereof, comprising:

- a) providing a recombinant parent antibody or antigen-binding portion thereof; that was obtained by selection in a phage-display system but whose activity cannot be further improved by mutagenesis in said phage-display system;
- b) selecting a selecting an amino acid residue within a complementarity

 determining region (CDR) for mutation other than H30, H31, H31B, H32, H33, H35,

 H50, H52, H52A, H53, H54, H56, H58, H95, H96, H97, H98, H101, L30, L31, L32,

 L34, L50, L52, L53, L55, L91, L92, L93, L94 and;

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c) individually mutating said selected contact or hypermutation position to at least two other amino acid residues to thereby create a panel of mutated antibodies, or antigen-binding portions thereof, and expressing said panel in a non-phage display evetore.

system;

d) evaluating the activity of the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof

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thereby identifying an activity enhancing amino acid residue;

e) evaluating the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof, for characteristic, until an antibody, or antigen-binding portion thereof, with an improved activity, relative to the parent antibody, or antigen-binding thereof, with an improved activity, relative to the parent antibody, or antigen-binding

portion thereof, is obtained.

Preferably, the other characteristic or property is selected from 1) preservation of

non-crossreactivity with other proteins or human tissues, 2) preservation of epitope recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 heterodimer preventing binding interference from free, soluble p40 and/or 3) to produce

recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 heterodimer preventing binding interference from free, soluble p40 and/or 3) to produce an antibody with a close to germline immunoglobulin sequence.

If a single mutagenesis is not sufficient to increase the affinity of the antibody

other residues may be included in the mutagenesis. Therefore, in another embodiment, the invention provides a method for improving the activity of an antibody, or antigen-

binding portion thereof, comprising:

a) providing a parent antibody or antigen-binding portion thereof that was

obtained by mutagenesis in said phage-display system;

b) selecting an amino acid residue within a complementarity determining region (CDR) for mutation other than H30, H31, H31B, H32, H35, H35, H50, H52, H52A,

c) individually mutating said selected position to at least two other amino acid thereof and expression in a non-phage display system;

2/L/98/00/4θ ZL/20/00/4θ ZL/20/00/20/00/20/20/0

d) evaluating the activity of the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof thereby identifying an activity enhancing amino acid residue;

L52, L53, L55, L91, L92, L93, L94;
g) combining, in the parent antibody, or antigen-binding portion thereof, at least two individual activity enhancing amino acid residues shown to have improved activity, to form combination antibodies, or antigen-binding portions thereof; and

h) evaluating the activity and other property or characteristic of the combination antibodies, or antigen-binding portions thereof with two activity enhancing amino acid residues, relative to the parent antibody or antigen-binding portion thereof, with an improved activity, relative to the parent antibody, or antigen-binding portion thereof, is obtained.
5 to the parent antibody, or antigen-binding portion thereof, is obtained.
Preferably, the other characteristic or property is selected from 1) preservation of preferably, the other characteristic or property is selected from 1) preservation of

non-crossreactivity with other proteins or human tissues, 2) preservation of epitope recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 heterodimer preventing binding interference from free, soluble p40 and/or 3) to produce an antibody with a close to germline immunoglobulin sequence.

The preferred attempts to identify activity enhancing amino acids focussing on

the preferred selective mutagenesis positions, contact or hypermutation positions described may be exhausted, or additional improvements may be required, and it is important to retain other properties or characteristics of the antibody.

Therefore, in another embodiment, the invention provides a method for

improving the activity of an antibody, or antigen-binding portion thereof, without affecting other characteristics, comprising:

a) providing a parent antibody or antigen-binding portion thereof;
b) selecting an amino acid residue within a complementarity determining region
(CDR) for mutation other than H30, H31, H31B, H32, H33, H35, H50, H52A,

T22' T31' T35' T34' H26' H32' H32' H36' H31' H38' H101' T30' T31' T35' T34' T20' T25' T23'

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c) individually mutating said selected position to at least two other amino acid residues to thereby create a panel of mutated antibodies, or antigen-binding portions

thereof;

d) evaluating the activity of the panel of mutated antibodies. or antigen-binding

portions thereof, relative to the parent antibody or antigen-binding portion thereof thereby identifying an activity enhancing amino acid residue;

e) evaluating the panel of mutated antibodies, or antigen-binding portions

thereof, relative to the parent antibody or antigen-binding portion thereof, for changes in at least one other property or characteristic until an antibody, or antigen-binding portion thereof, with an improved activity and retained other property or characteristic, relative

to the parent antibody, or antigen-binding portion thereof, is obtained.

Preferably, the other characteristic or property is selected from 1) preservation of

non-crossreactivity with other proteins or human tissues, 2) preservation of epitope recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 heterodimer preventing binding interference from free, soluble p40 and/or 3) to produce

an antibody with a close to germline immunoglobulin sequence

If mutagenesis of a single residue is not sufficient other residues can be included;

therefore, in another embodiment, the invention provides a method for improving the

activity of an antibody, or antigen-binding portion thereof, comprising:

a) providing a parent antibody or antigen-binding portion thereof;

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b) selecting an amino acid residue within a complementarity determining region (CDR) for mutation other than H30, H31, H31B, H32, H33, H35, H50, H52, H52A, H53, H54, H56, H58, H95, H97, H98, H101, L30, L31, L32, L34, L50, L52, L53,

L55, L91, L92, L93, L94 and L96;

c) individually mutating said selected position to at least two other amino acid residues to thereby create a panel of mutated antibodies, or antigen-binding portions

thereof;

d) evaluating the activity of the panel of mutated antibodies, or antigen-bindi

d) evaluating the activity of the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof, thereby identifying an activity enhancing amino acid residue;

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c.) evaluating the panel of mutated antibodies or antigen-binding portions thereof, relative to the parent antibody or antigen-portion thereof, for changes in at least

- one other characteristic or property;

 e) repeating steps b) through e) for at least one other CDR position which is neither the position selected under b) nor a position at H30, H31, H31B, H32, H33, H35
- neither the position selected under b) nor a position at H30, H31, H31B, H32, H33, H35, H50, H52, H52, H53, H54, H56, H56, H96, H97, H98, H101, L30, L31, L32, L34, L50, L52, L53, L54, L92, L93, L94, and L96;
- f) combining, in the parent antibody, or antigen-binding portion thereof, at least two individual activity enhancing amino acid residues shown to have improved activity
 and not affecting at least one other property or characteristic, to form combination
- antibodies, or antigen-binding portions thereof; and
 g) evaluating the activity and the retention of at least one other property or characteristic of the combination antibodies, or antiven-binding portions thereof with
- characteristic of the combination antibodies, or antigen-binding portions thereof with two activity enhancing amino acid residues, relative to the parent antibody or antigen-binding portion thereof until an antibody, or antigen-binding portion thereof, with an improved activity and at least one retained other property or characteristic, relative to the parent antibody, or antigen-binding portion thereof, is obtained.
- Mutagenesis of the preferred selective mutagenesis position, contact and hypermutation residues may not have increased the affinity of the antibody sufficiently, and mutagenesis and the phage display method (or related ribosome display method) may no longer be useful and at least one other characteristic or property of the antibody should be tetained
- should be retained.

 Therefore, in another embodiment the invention provides a method to improve the affinity of an antibody or antigen-binding portion thereof, comprising:
- a) providing a parent antibody or antigen-binding portion thereof that was obtained by selection in a phage-display system but whose activity cannot be further

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- L55, L91, L92, L94, and L96;

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c) individually mutating said selected position to at least two other amino acid residues to thereby create a panel of mutated antibodies, or antigen-binding portions thereof and expression in a non-phage display system;

- d) evaluating the activity of the parent antibody or antigen-binding portion thereof

 portions thereof, relative to the parent antibody or antigen-binding portion thereof
- thereby identifying an activity enhancing amino acid residue;

 e) evaluating the panel of mutated antibodies, or antigen-binding portions
- thereof, relative to the parent antibody or antigen-binding portion thereof, for changes in at least one other property or characteristic until an antibody, or antigen-binding portion thereof, with an improved activity, relative to the parent antibody, or antigen-binding

portion thereof, is obtained.

Preferably, the other characteristic or property is selected from 1) preservation of

non-crossreactivity with other proteins or human tissues, 2) preservation of epitope recognition, i.e. recognizing p40 epitope preferably in the context of the p70 p40/p35 heterodimer preventing binding interference from free, soluble p40 and/or 3) to produce

an antibody with a close to germline immunoglobulin sequence

If mutagenesis of a single residue is not sufficient other residues can be included;

therefore, in another embodiment, the invention provides a method for improving the activity of an antibody, or antigen-binding portion thereof, comprising:

a) providing a parent antibody or antigen-binding portion thereof that was obtained by selection in a phage-display system but whose activity cannot be further improved by mutscenesis in sold phage-display system.

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improved by mutagenesis in said phage-display system;

b) selecting an amino acid residue within a complementarity determining region

(GDP) for mytasien ether then 1130, 1131, 1132, 1132, 1131, 1132, 1133, 113

c) individually mutating said selected position to at least two other amino acid residues to thereby create a panel of mutated antibodies, or antigen-binding portions thereof and expression in a non-phage display system;

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d) evaluating the activity and retention of at least one other property or characteristic of the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof, thereby identifying an activity enhancing amino acid residue:

- e) repeating steps b) through d) for at least one other CDR position which is neither the position selected under b) nor a position at H30, H31, H31, H31, H32, H33, H35, H30, H52, L53, L53, L51, L93, L94, B06, H97, H98, H101, L30, L31, L32, L34, L50, L52, L53, L53, L91, L92, L93, L94 and L96;
- f) combining, in the parent antibody, or antigen-binding portion thereof, at least two individual activity enhancing amino acid residues shown to have improved activity and not to affect at least one other property or characteristic, to form combination antibodies, or antigen-binding portions thereof; and
- g) evaluating the activity and retention of at least one property or characteristic of the combination antibodies, or antigen-binding portions thereof with two activity characteristic or property, relative to the parent antibody, or antigen-binding portion thereof, with an improved activity and at least one other retained characteristic or property, relative to the parent antibody, or antigen-binding portion thereof, is obtained.

20 V Expression of Antibodies

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An antibody, or antibody portion, of the invention can be prepared by recombinant expression of immunoglobulin light and heavy chain genes in a host cell.

To express an antibody recombinantly, a host cell is transfected with one or more recombinant expression vectors carrying DNA fragments encoding the immunoglobulin in the host cell and, preferably, secreted into the medium in which the host cells are in the host cell and, preferably, secreted into the medium in which the host cells are cultured, from which medium the antibody heavy and light chain genes, into host cells, such as those described in Sambrook, Fritsch and Maniatis (eds), into host cells, such as those described in Sambrook, Fritsch and Maniatis (eds), into host cells, such as those described in Sambrook, Fritsch and Maniatis (eds), into host cells, such as those described in Sambrook, Fritsch and Maniatis (eds), into host cells, such as those described in Sambrook, Fritsch and Maniatis (eds), into host cells, such as those described in Sambrook, Fritsch and Maniatis (eds), into host cells, such as those described in Sambrook, Fritsch and Maniatis (eds), into host cells, such as those described in Sambrook, Fritsch and Maniatis (eds), into host cells, such as those described in Sambrook, Fritsch and Maniatis (eds), into host cells, such as those described in Sambrook, Fritsch and Maniatis (eds), into host cells, such as those described in Sambrook, Fritsch and Maniatis (eds), into host cells, such as those described in Sambrook, Fritsch and Maniatis (eds), into host cells, such as those described in Sambrook, Fritsch and Maniatis (eds), into host cells, such as those described in Sambrook, Fritsch and Proposition when the host cells, into host c

(1989), Ausubel, F.M. et al. (eds.) Current Protocols in Molecular Biology, Greene Publishing Associates, (1989) and in U.S. Patent No. 4,816.397 by Boss et al.

To obtain a DNA fragment encoding the heavy chain variable region of loe 9 wt

or a loe 9 wt-related antibody, antibodies specific for human IL-12 were screened from human libraries and mutated, as described in section II. Once DNA fragments encoding loe 9 wt or loe 9 wt-related VH and VL segments are obtained, mutagenesis of these sequences is carried out by standard methods, such as PCR site directed mutagenesis (PCR-mediated mutagenesis in which the mutated nucleotides are incorporated into the PCR primers such that the PCR product contains the mutations) or other site-directed mutagenesis methods. Human IL-12 antibodies that displayed a level of activity and binding specificity/affinity that was desirable, for example 1695, were further

binding specificity/affinity that was desirable, for example 1695, were further manipulated by standard recombinant DNA techniques, for example to convert the variable region genes to full-length antibody chain genes, to Fab fragment genes or to a scFv gene. In these manipulations, a VL- or VH-encoding DNA fragment is operatively linked to another DNA fragment encoding another protein, such as an antibody constant region or a flexible linker. The term "operatively linked", as used in this context, is

region or a flexible linker. The term "operatively linked", as used in this context, is intended to mean that the two DNA fragments are joined such that the amino acid sequences encoded by the two DNA fragments remain in-frame.

The isolated DNA encoding the VH region can be converted to a full-length

nolecule encoding heavy chain constant regions (CHI, CH2 and CH3). The sequences of human heavy chain constant regions (CHI, CH2 and CH3). The sequences of human heavy chain constant region genes are known in the art (see e.g., Kabat, E.A., et al. (1991) Sequences of Proteins of Immunological Interest, Fifth Edition, U.S.

The heavy chain constant region can be obtained by standard PCR amplification. The heavy chain constant region can be obtained by standard PCR amplification. IgD constant region and any allotypic variant therein as described in Kabat (, Kabat, IgD constant region and any allotypic variant therein as described in Kabat (, Kabat, IgD constant region and any allotypic variant therein as described in Kabat (, Kabat, IgD constant region and human Services, NIH Publication No. 91-3242), but most Department of Health and Human Services, MIH Publication No. 91-3242), but most preferably is an IgG1 or IgG4 constant region. For a Fab fragment heavy chain gene, the VH-encoding DNA can be operatively linked to another DNA molecule encoding the VH-encoding DNA can be operatively linked to another DNA molecule encoding

only the heavy chain CH1 constant region.

The isolated DNA encoding the VL region can be converted to a full-length light chain gene (as well as a Fab light chain gene) by operatively linking the VL-encoding DNA to another DNA molecule encoding the light chain constant region, CL. The sequences of human light chain constant region genes are known in the art (see e.g., Edition, U.S. Department of Health and Human Services, NIH Publication No. 91-3242) and DNA fragments encompassing these regions can be obtained by standard PCR amplification. The light chain constant region can be a kappa or lambda constant region, but most preferably is a lambda constant region.

To create a seFv gene, the VH- and VL-encoding DNA fragments are operatively or the operative of the VH- and VL-encoding DNA fragments are operatively or the operative of the VH- and VL-encoding DNA fragments are operatively or the operative of the VH- and VL-encoding DNA fragments are operatively or the operative of the VH- and VL-encoding DNA fragments are operatively to the operative of the

linked to another fragment encoding a flexible linker, e.g., encoding the amino acid sequence (Gly₄-Set)₃, such that the VH and VL sequences can be expressed as a contiguous single-chain protein, with the VL and VH regions joined by the flexible linker (see e.g., Bird et al. (1988) Science 242:423-426; Huston et al. (1988) Proc. Natl. Acad. Sci. USA 85:5879-5883; McCafferty et al., Nature (1990) 348:552-554).

partial or full-length light and heavy chains, obtained as described above, are inserted into expression vectors such that the genes are operatively linked to transcriptional and translational control sequences. In this context, the term "operatively linked" is intended translational control sequences within the vector serve their intended function of translational control sequences within the vector serve their intended function of and expression control sequences are chosen to be compatible with the expression vector cell used. The antibody light chain gene and the antibody heavy chain gene can be inserted into separate vector or, more typically, both genes are inserted into the same expression vector. The antibody genes are inserted into the same standard methods (e.g., ligation of complementary restriction sites on the antibody gene standard methods (e.g., ligation of complementary restriction sites on the antibody gene

approach to converting the J695 or J695-related VH and VL sequences to full-length antibody genes is to insert them into expression vectors already encoding heavy chain

vector may already carry antibody constant region sequences. For example, one

insertion of the J695 or J695-related light or heavy chain sequences, the expression

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constant and light chain constant regions, respectively, such that the VH segment is operatively linked to the CH segment(s) within the vector and the VL segment is operatively linked to the CL segment within the vector. Additionally or alternatively, the recombinant expression vector can encode a signal peptide that facilitates secretion of the antibody chain from a host cell. The antibody chain gene can be cloned into the vector such that the signal peptide is linked in-frame to the amino terminus of the vector such that the signal peptide can be an immunoglobulin signal peptide or a antibody chain gene. The signal peptide can be an immunoglobulin signal peptide or a

In addition to the antibody chain genes, the recombinant expression vectors of

heterologous signal peptide (i.e., a signal peptide from a non-immunoglobulin protein).

No. 4,968,615 by Schaffner et al., U.S. Patent No.5, 464,758 by Bujard et al. and U.S. Patent No. 5,168,062 by Stinski, U.S. Patent No. 4,510,245 by Bell et al. and U.S. Patent further description of viral regulatory elements, and sequences thereof, see e.g., U.S. adenovirus, (e.g., the adenovirus major late promoter (AdMLP)) and polyoma. For promoter/enhancer), Simian Virus 40 (SV40) (such as the SV40 promoter/enhancer), promoters and/or enhancers derived from cytomegalovirus (CMV) (such as the CMV elements that direct high levels of protein expression in mammalian cells, such as Preferred regulatory sequences for mammalian host cell expression include viral of the host cell to be transformed, the level of expression of protein desired, etc. including the selection of regulatory sequences may depend on such factors as the choice will be appreciated by those skilled in the art that the design of the expression vector, Technology: Methods in Enzymology 185, Academic Press, San Diego, CA (1990). It ۶ı regulatory sequences are described, for example, in Goeddel; Gene Expression signals) that control the transcription or translation of the antibody chain genes. Such promoters, enhancers and other expression control elements (e.g., polyadenylation chain genes in a host cell. The term "regulatory sequence" is intended to include the invention carry regulatory sequences that control the expression of the antibody

In addition to the antibody chain genes and regulatory sequences, the recombinant expression vectors of the invention may carry additional sequences, such as and selectable marker genes. The selectable marker gene facilitates selection of host cells into which the vector has been introduced (see e.g., U.S. Patents Nos. 4,399,216,

Patent No. 5,654,168 by Bujard et al.

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4,634,665 and 5,179,017, all by Axel et al.). For example, typically the selectable marker gene confers resistance to drugs, such as G418, hygromycin or methotrexate, on a host cell into which the vector has been introduced. Preferred selectable marker genes include the dihydrofolate reductase (DHFR) gene (for use in dhfr⁻ host cells with methotrexate selection/amplification) and the neo gene (for G418 selection).

For expression of the light and heavy chains, the expression vector(s) encoding

the heavy and light chains is transfected into a host cell by standard techniques. The various forms of the term "transfected into a host cell by standard techniques. The techniques commonly used for the introduction of exogenous DNA into a prokaryotic or dextran transfection and the like. Although it is theoretically possible to express the antibodies of the invention in either prokaryotic or eukaryotic host cells, expression of antibodies in eukaryotic cells, and most preferably mammalian host cells, is the most preferred because such eukaryotic cells, and in particular mammalian cells, is the most preferred because such eukaryotic cells, and in particular mammalian cells, are more likely than prokaryotic cells to assemble and secrete a properly folded and immunologically active antibody. Preferred mammalian host cells for expressing the recombinant antibodies of the invention include Chinese Hamster Ovary (CHO cells) (including dhfr- CHO cells, described in Urlaub and Chasin, (1980) Proc. Mail. Acad. (including dhfr- CHO cells, described in Urlaub and Chasin, (1980) Proc. Mail. Acad.

Kaufman and P.A. Sharp (1982) *Mol. Biol.* 159:601-621), NS0 myeloma cells, COS cells and SP2 cells. When recombinant expression vectors encoding antibody genes are introduced into mammalian host cells, the antibodies are produced by culturing the host cells for a period of time sufficient to allow for expression of the antibody in the host cells or, more preferably, secretion of the antibody into the culture medium in which the cells or, more preferably, secretion of the antibody into the culture medium in which the

Sci. VSA $\overline{11}$:4216-4220, used with a DHFR selectable marker, e.g., as described in R.J.

standard protein purification methods.

Host cells can also be used to produce portions of intact antibodies, such as Fab

fragments or scFv molecules. It will be understood that variations on the above procedure are within the scope of the present invention. For example, it may be desirable to transfect a host cell with DNA encoding either the light chain or the heavy chain (but not both) of an antibody of this invention. Recombinant DNA technology may also be used to remove some or all of the DNA encoding either or both of the light

and heavy chains that is not necessary for binding to hIL-12. The molecules expressed from such truncated DNA molecules are also encompassed by the antibodies of the invention. In addition, bifunctional antibodies may be produced in which one heavy and one light chain are an antibody of the invention and the other heavy and light chain are specific for an antigen other than hIL-12 by crosslinking an antibody of the invention to a second antibody by standard chemical crosslinking methods.

In a preferred system for recombinant expression of an antibody, or antigen-binding portion thereof, of the invention, a recombinant expression vector encoding both the antibody heavy chain and the antibody light chain is introduced into dhfr- CHO cells by calcium phosphate-mediated transfection. Within the recombinant expression vector,

In view of the foregoing, another aspect of the invention pertains to nucleic acid, plants that express the antibody or antigen binding portion thereof, of the invention. Nucl. Acids Res. 20: 6287-6295). Plant cells can also be modified to create transgenic that is transgenic for human immunoglobulin genes (see e.g., Taylor, L.D. et al. (1992) binding portions thereof of the invention can be expressed in an animal (e.g., a mouse) the host cells and recover the antibody from the culture medium. Antibodies or antigenrecombinant expression vector, transfect the host cells, select for transformants, culture the culture medium. Standard molecular biology techniques are used to prepare the expression of the antibody heavy and light chains and intact antibody is recovered from selection/amplification. The selected transformant host cells are culture to allow for for selection of CHO cells that have been transfected with the vector using methotrexate the genes. The recombinant expression vector also carries a DHFR gene, which allows enhancer/AdMLP promoter regulatory element) to drive high levels of transcription of the like, such as a CMV enhancer/AdMLP promoter regulatory element or an SV40 enhancer/promoter regulatory elements (e.g., derived from SV40, CMV, adenovirus and the antibody heavy and light chain genes are each operatively linked to

vector and host cell compositions that can be used for recombinant expression of the antibodies and antibody portions of the invention. Preferably, the invention features isolated nucleic acids that encode CDRs of 1695, or the full heavy and/or light chain variable region of 1695. Accordingly, in one embodiment, the invention features an isolated nucleic acid encoding an antibody heavy chain variable region that encodes the lesson that encodes the solution that confirm that encodes the solution that confirm that encodes the solution that confirm that encodes the lesson chain CDR3 comprising the amino acid sequence of SEQ ID NO: 25.

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Preferably, the nucleic acid encoding the antibody heavy chain variable region further encodes a 1695 heavy chain CDR2 which comprises the amino acid sequence of SEQ ID MO: 27. More preferably, the nucleic acid encoding the antibody heavy chain variable region further encodes a 1695 heavy chain CDR1 which comprises the amino acid sequence of SEQ ID MO: 29. Even more preferably, the isolated nucleic acid encodes an antibody heavy chain variable region comprising the amino acid sequence of SEQ ID MO: 31 (the full VH region of 1695).

an antibody light chain variable region that encodes the J695 light chain CDR3 comprising the amino acid sequence of SEQ ID NO: 26. Preferably, the nucleic acid encoding the antibody light chain variable region further encodes a J695 light chain the nucleic acid encoding the antibody light chain variable region further encodes a J695 light chain CDR1 which comprises the amino acid sequence of SEQ ID NO: 30. Even light chain CDR1 which comprises the amino acid sequence of SEQ ID NO: 30. Even light chain comprising the amino acid sequence of SEQ ID NO: 31. Even region comprising the amino acid sequence of SEQ ID NO: 32 (the full VL region of region comprising the amino acid sequence of SEQ ID NO: 32 (the full VL region of region comprising the amino acid sequence of SEQ ID NO: 32 (the full VL region of region comprising the amino acid sequence of SEQ ID NO: 32 (the full VL region of region comprising the amino acid sequence of SEQ ID NO: 32 (the full VL region of region comprising the amino acid sequence of SEQ ID NO: 32 (the full VL region of region comprising the amino acid sequence of SEQ ID NO: 32 (the full VL region of region comprising the amino acid sequence of SEQ ID NO: 32 (the full VL region of region comprising the amino acid sequence of SEQ ID NO: 32 (the full VL region of region comprising the amino acid sequence of SEQ ID NO: 32 (the full VL region of region comprising the amino acid sequence of SEQ ID NO: 32 (the full VL region of region comprising the amino acid sequence of SEQ ID NO: 32 (the full VL region of region comprising the amino acid sequence of SEQ ID NO: 32 (the full VL region of region comprising the acid sequence of SEQ ID NO: 32 (the full VL region of region comprising the acid sequence of SEQ ID NO: 32 (the full VL region of region comprising the acid sequence of SEQ ID NO: 32 (the full VL region of region comprising the acid sequence of SEQ ID NO: 32 (the full VL region of region comprising the acid sequence of SEQ ID NO: 32 (the full VL region of region comprising the acid sequenc

In other embodiments, the invention features an isolated nucleic acid encoding

region comprising the amino acid sequence of SEQ ID NO: 32 (the full VL region of J695).

The invention also provides recombinant expression vectors encoding both an

antibody heavy chain and an antibody light chain. For example, in one embodiment, the invention provides a recombinant expression vector encoding:

a) an antibody heavy chain having a variable region comprising the

amino acid sequence of SEQ ID NO: 31; and
b) an antibody light chain having a variable region comprising the amino

acid sequence of SEQ ID NO: 32.

The invention also provides host cells into which one or more of the recombinant

expression vectors of the invention have been introduced. Preferably, the host cell is a mammalian host cell, more preferably the host cell is a CHO cell, an NS0 cell or a COS cell. Still further the invention provides a method of synthesizing a recombinant human antibody of the invention in a suitable culture medium until a recombinant human antibody of the invention is synthesized. The method can further comprise isolating the recombinant human antibody from the culture

medium.

VI. Pharmaceutical Compositions and Pharmaceutical Administration

The antibodies and antibody-portions of the invention can be incorporated into pharmaceutical compositions suitable for administration to a subject. Typically, the and a pharmaceutical composition comprises an antibody or antibody portion of the invention and a pharmaceutically acceptable carrier. As used herein, "pharmaceutically acceptable carrier and all solvents, dispersion media, coatings, antibacterial and antifungal agents, isotonic and absorption delaying agents, and the like that are antifungal agents, isotonic and absorption delaying agents, and the like that are one or more of water, saline, phosphate buffered saline, dextrose, glycerol, ethanol and one or more of water, saline, phosphate buffered saline, dextrose, glycerol, ethanol and

one or more of water, saline, phosphate buffered saline, dextrose, glycerol, ethanol and the like, as well as combinations thereof. In many cases, it will be preferable to include isotonic agents, for example, sugars, polyalcohols such as mannitol, sorbitol, or sodium chloride in the composition. Pharmaceutically acceptable carriers may further comprise minor amounts of auxiliary substances such as wetting or emulsifying agents, preservatives or buffers, which enhance the shelf life or effectiveness of the antibody or

antibody portion.

The antibodies and antibody-portions of the invention can be incorporated into a

pharmaceutical composition suitable for parenteral administration. Preferably, the antibody or antibody-portions will be prepared as an injectable solution containing 0.1-150 mg/ml antibody. The injectable solution can be composed of either a liquid or lyophilized dosage form in a flint or amber vial, ampule or pre-filled syringe. The buffer can be L-histidine (1-50 mM), optimally 5-10mM, at pH 5.0 to 7.0 (optimally pH 6.0). Other suitable buffers include but are not limited to, sodium succinate, sodium citrate,

dosage form). Cryoprotectants can be included for a lyophilized dosage form, principally 0-10% sucrose (optimally 0.5-1.0%). Other suitable cryoprotectants include treahalose and lactose. Bulking agents can be included for a lyophilized dosage form, principally 1-10% mannitol (optimally 2-4%). Stabilizers can be used in both liquid and lyophilized dosage forms, principally 1-50 mM L-Methionine (optimally 5-10 mM).

Other suitable bulking agents include glycine, arginine, can be included as 0-0.05%

sodium phosphate or potassium phosphate. Sodium chloride can be used to modify the toxicity of the solution at a concentration of 0-300 mM (optimally 150 mM for a liquid

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polysorbate-80 (optimally 0.005-0.01%). Additional surfactants include but are not limited to polysorbate 20 and BRIJ surfactants.

In a preferred embodiment, the pharmaceutical composition includes the antibody at a dosage of about 0.01mg/kg - 10 mg/kg. More preferred dosages of the antibody include Img/kg administered every other week, or 0.3 mg/kg administered weekly.

The compositions of this invention may be in a variety of forms. These include, for example, liquid, semi-solid and solid dosage forms, such as liquid solutions (e.g., injectable and infusible solutions), dispersions or suspensions, tablets, pills, powders, liposomes and suppositories. The preferred form depends on the intended mode of administration and therapeutic application. Typical preferred compositions are in the

liposomes and suppositories. The preferred form depends on the intended mode of administration and therapeutic application. Typical preferred compositions are in the form of injectable or infusible solutions, such as compositions similar to those used for passive immunization of humans with other antibodies. The preferred mode of administration is parenteral (e.g., intravenous, subcutaneous, intraperitoneal, intramuscular). In a preferred embodiment, the antibody is administered by intravenous infusion or injection. In another preferred embodiment, the antibody is administered by intravenous infusion or injection. In another preferred embodiment, the antibody is administered by intravenous infusion or injection. In another preferred embodiment, the antibody is administered by intravenous infusion or injection.

intramuscular or subcutaneous injection.

Therapeutic compositions typically must be sterile and stable under the conditions

of manufacture and storage. The composition can be formulated as a solution,

microemulsion, dispersion, liposome, or other ordered structure suitable to high drug concentration. Sterile injectable solutions can be prepared by incorporating the active compound (i.e., antibody or antibody portion) in the required amount in an appropriate solvent with one or a combination of ingredients enumerated above, as required, followed by filtered sterilization. Generally, dispersions are prepared by incorporating the active other ingredients from those enumerated above. In the case of sterile, lyophilized powders other ingredients from those enumerated above. In the preferred methods of preparation are vacuum drying and spray-drying that yields a powder of the active ingredient plus any additional desired ingredient from a previously sterile-filtered solution thereof. The active ingredient plus any broper fluidity of a solution can be maintained, for example, by the use of a coating such as lecithin, by the maintenance of the required particle size in the case of dispersion and by the use of surfactants. Prolonged absorption of injectable compositions can be brought by the use of surfactants. Prolonged absorption of injectable compositions can be brought

about by including in the composition an agent that delays absorption, for example, monostearate salts and gelatin.

The antibodies and antibody-portions of the present invention can be administered by a variety of methods known in the art, although for many therapeutic applications, the preferred route/mode of administration is subcutaneous injection, intravenous injection or

infusion. As will be appreciated by the skilled artisan, the route and/or mode of administration will vary depending upon the desired results. In certain embodiments, the active compound may be prepared with a carrier that will protect the compound against rapid release, such as a controlled release formulation, including implants, transdermal polymers can be used, such as ethylene vinyl acetate, polyanhydrides, polyglycolic acid, polymers can be used, such as ethylene vinyl acetate, polyanhydrides, polyglycolic acid, collagen, polyorthoesters, and polylactic acid. Many methods for the preparation of such formulations are patented or generally known to those skilled in the art. See, e.g., Sustained and Controlled Release Drug Delivery Systems, J.R. Robinson, ed., Marcel Dekker, Inc., New York, 1978.

In certain embodiments, an antibody or antibody portion of the invention may be orally administered, for example, with an inert diluent or an assimilable edible carrier.

The compound (and other ingredients, if desired) may also be enclosed in a hard or soft shell gelatin capsule, compressed into tablets, or incorporated directly into the subject's diet. For oral therapeutic administration, the compounds may be incorporated with excipients and used in the form of ingestible tablets, buccal tablets, troches, capsules, excipients and used in the form of ingestible tablets, buccal tablets, troches, capsules, elixirs, suspensions, syrups, wafers, and the like. To administer a compound of the invention by other than parenteral administration, it may be necessary to coat the compound with, or co-administer the compound with, a material to prevent its

Supplementary active compounds can also be incorporated into the compositions. In certain embodiments, an antibody or antibody portion of the invention is coformulated with and/or coadministered with one or more additional therapeutic agents that are useful for treating disorders in which IL-12 activity is detrimental. For example, an anti-hIL-12 antibody or antibody portion of the invention may be coformulated and/or coadministered with one or more additional antibodies that bind other targets (e.g., antibodies that bind other cytokines or that bind cell surface

inactivation.

molecules). Furthermore, one or more antibodies of the invention may be used in combination with two or more of the foregoing therapeutic agents. Such combination with two or more of the foregoing therapeutic agents. Such combination agents, thus avoiding possible toxicities or complications associated with the various monotherapies. It will be appreciated by the skilled practitioner that when the antibodies of the invention are used as part of a combination therapy, a lower dosage of antibody of the invention are used as part of a combination therapy, a lower dosage of antibody aynergistic therapeutic effect may be achieved through the use of combination therapy which, in turn, permits use of a lower dose of the antibody to achieve the desired therapuetic effect).

Interleukin 12 plays a critical role in the pathology associated with a variety of

pemphigus vulgaris, pemphigus foliaceus, pemphigoid, linear IgA disease, autoimmune atheromatous disease/arteriosclerosis, atopic allergy, autoimmune bullous disease, chlamydia, yersinia and salmonella associated arthropathy, spondyloarthopathy, 30 disease, psoriatic arthropathy, ulcerative colitic arthropathy, enteropathic synovitis, syndrome, alopecia, alopecia areata, seronegative arthopathy, arthropathy, Reiter's polyglandular deficiency type II, Schmidt's syndrome, adult (acute) respiratory distress myocardial infarction, Addison's disease, sporadic, polyglandular deficiency type I and stroke, primary biliary cirrhosis, hemolytic anemia, malignancies, heart failure, transverse myelitis, Huntington's chorea, Parkinson's disease, Alzheimer's disease, infectious diseases, parasitic diseases, acquired immunodeficiency syndrome, acute hepatitis, uveitis, septic shock, toxic shock syndrome, sepsis syndrome, cachexia, Henoch-Schoenlein purpurea, microscopic vasculitis of the kidneys, chronic active disease, nephrotic syndrome, chronic fatigue syndrome, Wegener's granulomatosis, atherosclerosis, disseminated intravascular coagulation, Kawasaki's disease, Grave's acute or chronic immune disease associated with organ transplantation, sarcoidosis, scleroderma, atopic dermatitis, graft versus host disease, organ transplant rejection, dependent diabetes mellitus, thyroiditis, asthma, allergic diseases, psoriasis, dermatitis erythematosus, Crohn's disease, ulcerative colitis, inflammatory bowel disease, insulin arthritis, psoriatic arthritis, reactive arthritis, spondyloarthropathy, systemic lupus not limited to, rheumatoid arthritis, osteoarthritis, juvenile chronic arthritis, Lyme diseases involving immune and inflammatory elements. These diseases include, but are

myxoedema, phacogenic uveitis, primary vasculitis and vitiligo. The human antibodies, hypothyroidism (Hashimoto's disease), atrophic autoimmune hypothyroidism, primary thrombocytopenia; autoimmune thyroid disease, hyperthyroidism, goitrous autoimmune Takayasu's disease/arteritis, autoimmune thrombocytopenia, idiopathic nodosa, acute rheumatic fever, rheumatoid spondylitis, Still's disease, systemic sclerosis, tissue disease, Goodpasture's syndrome, pulmonary manifestation of polyarteritis mellitus, sympathetic ophthalmia, pulmonary hypertension secondary to connective sperm autoimmunity, multiple sclerosis (all subtypes), insulin-dependent diabetes kidneys, lyme disease, discoid lupus erythematosus, male infertility idiopathic or MOS, neutropenia, renal disease NOS, glomerulonephritides, microscopic vasulitis of the osteoarthrosis, primary sclerosing cholangitis, idiopathic leucopenia, autoimmune transplantation, chronic immune disease associated with organ transplantation, acanthosis nigricans, hypoparathyroidism, acute immune disease associated with organ hepatitis), autoimmune mediated hypoglycemia, type B insulin resistance with autoimmune or lupoid hepatitis), type-2 autoimmune hepatitis (anti-LKM antibody gouty arthritis, autoimmune hepatitis, type-1 autoimmune hepatitis (classical pneumonia, lymphocytic infiltrative lung disease, postinfectious interstitial lung disease, interstitial lung disease, radiation fibrosis, bronchiolitis obliterans, chronic eosinophilic vasculitic diffuse lung disease, haemosiderosis associated lung disease, drug-induced disease associated lung disease, ankylosing spondylitis associated lung disease, associated lung disease, dermatomyositis/polymyositis associated lung disease, Sjögren's rheumatoid arthritis associated interstitial lung disease, systemic lupus erythematosus disease associated lung disease, systemic seletosis associated interstitial lung disease, connective tissue disease associated interstitial lung disease, mixed connective tissue fibrosing alveolitis, post-inflammatory interstitial lung disease, interstitial pneumonitis, infertility, ovarian failure, premature ovarian failure, fibrotic lung disease, cryptogenic (common variable hypogammaglobulinaemia), dilated cardiomyopathy, female Immunodeficiency Related Diseases, Hepatitis C, common varied immunodeficiency autoimmune hepatitis, Acquired Immunodeficiency Disease Syndrome, Acquired mucocutaneous candidiasis, giant cell arteritis, primary sclerosing hepatitis, cryptogenic anaemia, juvenile pernicious anaemia, myalgic encephalitis/Royal Free Disease, chronic haemolytic anaemia. Coombs positive haemolytic anaemia, acquired pernicious

and antibody portions of the invention can be used to treat autoimmune diseases, in particular those associated with inflammation, including, rheumatoid spondylitis, allerey, autoimmune diabetes, autoimmune uveitis.

allergy, autoimmune diabetes, autoimmune uveitis.

Preferably, the antibodies of the invention or antigen-binding portions thereof, are used to treat rheumatoid arthritis, Crohn's disease, multiple selerosis, insulin

dependent diabetes mellitus and psoriasis, as described in more detail in section VII.

A human antibody, or antibody portion, of the invention also can be administered

with one or more additional therapeutic agents useful in the treatment of autoimmune and inflammatory diseases.

Antibodies of the invention, or antigen binding portions thereof can be used

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alone or in combination to treat such diseases. It should be understood that the antibodies of the invention or antigen binding portion thereof can be used alone or in combination with an additional agent, e.g., a therapeutic agent, said additional agent the being selected by the skilled artisan for its intended purpose. For example, the additional agent can be a therapeutic agent art-recognized as being useful to treat the disease or condition being treated by the antibody of the present invention. The additional agent also can be an agent which imparts a beneficial attribute to the therapeutic composition e.g., an agent which effects the viscosity of the composition.

therapeutic composition e.g., an agent which effects the viscosity of the composition.

It should further be understood that the combinations which are to be included within this invention are those combinations useful for their intended purpose. The agents set forth below are illustrative for purposes and not intended to be limited. The combinations which are part of this invention can be the antibodies of the present intention and at least one additional agents selected from the lists below. The

combinations which are part of this invention can be the antibodies of the present invention and at least one additional agent selected from the lists below. The combination can also include more than one additional agent, e.g., two or three additional agents if the combination is such that the formed composition can perform its intended function.

to as NSAIDS which include drugs like ibuprofen. Other preferred combinations are corticosteroids including prednisolone; the well known side-effects of steroid use can be reduced or even eliminated by tapering the steroid dose required when treating patients in combination with the anti-IL-12 antibodies of this invention. Non-limiting examples of therapeutic agents for rheumatoid arthritis with which an antibody, or antibody

Preferred combinations are non-steroidal anti-inflammatory drug(s) also referred

portion, of the invention can be combined include the following: cytokine suppressive anti-inflammatory drug(s) (CSAIDs); antibodies to or antagonists of other human cytokines or growth factors. for example, TMF, LT, IL-1, IL-2, IL-6. IL-7, IL-8, IL-15, IL-18, EMAP-II, GM-CSF, FGF, and PDGF. Antibodies of the invention, or antigen binding portions thereof, can be combined with antibodies to cell surface molecules such as CD2, CD3, CD4, CD8, CD25, CD28, CD30, CD40, CD45, CD69, CD80 (B7.1), CD86 (B7.2), CD90, or their ligands including CD154 (gp39 or CD40L). Preferred combinations of therapeutic agents may interfere at different points in the autoimmune and subsequent inflammatory cascade; preferred examples include TWF antagonists like chimeric, humanized or human TWF antibodies, D2E7, (U.S. application antagonists like chimeric, humanized or human TWF antibodies, D2E7, (U.S. application

The antibodies of the invention, or antigen binding portions thereof, may also be including antibodies, soluble receptors or antagonistic ligands. include antagonists of the co-stimulatory pathway CD80 (B7.1) or CD86 (B7.2) combination are non-depleting anti-CD4 inhibitors. Yet other preferred combinations and a combination of antagonists to both may be most effective. Yet another preferred proteins. It has been shown that IL-12 and IL-18 have overlapping but distinct functions 18 antagonists including IL-18 antibodies or soluble IL-18 receptors, or IL-18 binding. parallel to, dependent on or in concert with IL-12 function; especially preferred are ILpreferred combination are other key players of the autoimmune response which may act Interleukin 11, anti-P7s and p-selectin glycoprotein ligand (PSGL). Yet another IRA etc.) may be effective for the same reason. Other preferred combinations include IL-1 inhibitors (e.g., Interleukin-1-converting enzyme inhibitors, such as Vx740, orIL-13 receptor (sIL-13), and also TNF α converting enzyme (TACE) inhibitors; similarly derivatives thereof, (p75TNFR1gG (Enbrelim) or p55TNFR1gG (Lenercept), soluble IL-TNF antibody fragments (e.g., CDP870), and soluble p55 or p75 TNF receptors, serial number 08/599,226 filed February 9, 1996), cA2 (Remicade¹³¹). CDP 571, anti-

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combined with agents, such as methotrexate, 6-MP, azathioprine sulphasalazine, mesalazine, olsalazine chloroquinine/hydroxychloroquine, pencillamine, aurothiomalate (intramuscular and oral), azathioprine, cochicine, corticosteroids (oral, inhaled and local injection), beta-2 adrenoreceptor agonists (salbutamol, terbutaline, salmeteral), xanthines (theophylline, aminophylline), cromoglycate, nedocromil, ketotifen, ipratropium and oxitropium, cyclosporin, FK506, rapamycin, mycophenolate mofetil, leflunomide,

MSAIDs. for example, ibuprofen, corticosteroids such as prednisolone, phosphodiesterase inhibitors, adensosine agonists. antithrombotic agents, complement inhibitors, adrenergic agents, agents which interfere with signalling by proinflammatory cytokines such as TMFα or IL-1 (e.g. IRAK, MIK, IKK, p38 or MAP kinase inhibitors).

IL-1β converting enzyme inhibitors (e.g., Vx740), anti-P7s, p-selectin glycoprotein auch as kinase inhibitors, metalloproteinase inhibitors, aulfasalazine, azathioprine, 6-such as kinase inhibitors, metalloproteinase inhibitors, soluble cytokine receptors and derivatives thereof (e.g. soluble p55 or p75 TMF receptors and the derivatives py5TMFRIgG (EnbrelTM) and p55TMFRIgG (Lenercept), sIL-1RI, sIL-1RII, sIL-6R, p75TMFRIgG (EnbrelTM)) and antiinflammatory cytokines (e.g. IL-4, IL-10, IL-11, IL-13 and TGFβ). Preferred combinations include methotrexate or leflunomide and in moderate or severe rheumatoid arthritis cases, cyclosporine.

Non-limiting examples of therapeutic agents for inflammatory bowel disease

IL-1 β converting enzyme inhibitors (e.g., $\nabla x 740$), anti-P7s, p-selectin glycoprotein cytokines such as TNFa or IL-1 (e.g. IRAK, NIK, IKK, p38 or MAP kinase inhibitors), inhibitors, adrenergic agents, agents which interfere with signalling by proinflammatory phosphodiesterase inhibitors, adenosine agonists, antithrombotic agents, complement leflunomide, NSAIDs, for example, ibuprofen, corticosteroids such as prednisolone, such as methotrexate, cyclosporin, FK506, rapamycin, mycophenolate mofetil, of the invention, or antigen binding portions thereof, may also be combined with agents, CD8, CD25, CD28, CD30, CD40, CD45, CD69, CD90 or their ligands. The antibodies can be combined with antibodies to cell surface molecules such as CD2, CD3, CD4, CSF, FGF, and PDGF. Antibodies of the invention, or antigen binding portions thereof, example, TNF, LT, IL-1, IL-2, IL-6, IL-7, IL-8, IL-15, IL-16, IL-18, EMAP-II, GMcompounds; antibodies to or antagonists of other human cytokines or growth factors, for monoclonal antibodies; growth factors; elastase inhibitors; pyridinyl-imidazole inhibitors; IL-1 receptor antagonists; anti-IL-1 monoclonal antibodies; anti-IL-6 lipoxygenase inhibitors; mesalamine; olsalazine; balsalazide; antioxidants; thromboxane sulfasalazine; aminosalicylates; 6-mercaptopurine; azathioprine; metronidazole; the following: budenoside; epidermal growth factor; corticosteroids; cyclosporin, with which an antibody, or antibody portion, of the invention can be combined include

ligand (PSGL), TNFα converting enzyme inhibitors, T-cell signalling inhibitors such as kinase inhibitors, metalloproteinase inhibitors, sulfasalazine, azathioprine, 6-mercaptopurines, angiotensin converting enzyme inhibitors, soluble cytokine receptors and derivatives thereof (e.g. soluble p55 or p75 TNF receptors, sIL-1RII, sIL-8R, soluble IL-13 receptor (sIL-13)) and antiinflammatory cytokines (e.g. IL-4, IL-10,

IL-11, IL-13 and TGF\$).

Preferred examples of therapeutic agents for Crohn's disease in which an antibody or an antigen binding portion can be combined include the following: TWF

antagonists, for example, anti-TNF antibodies, D2E7 (U.S. application serial number

fragments (e.g., CDP870), TMFR-1g constructs(p75TMFR1gG (Enbre173) and fragments (e.g., CDP870), TMFR-1g constructs(p75TMFR1gG (Enbre173) and p55TMFR1gG (Lenercept)), anti-P7s, p-selectin glycoprotein ligand (PSGL), soluble ILbinding portions thereof, can be combined with corricosteroids, for example, budenoside and dexamethasone. Antibodies of the invention or antigen binding portions thereof, and agents auch as sulfasalazine, 5-aminosalicylic acid and olsalazine, and agents which interfere with synthesis or action of proinflammatory olsalazine, and agents which interfere with synthesis or action of proinflammatory cytokines such as IL-1, for example, IL-1\$ converting enzyme inhibitors (e.g., Vx740) and IL-1ra. Antibodies of the invention or antigen binding portion thereof may also be used with T cell signaling inhibitors, for example, tyrosine kinase inhibitors 6-

mercaptopurines. Antibodies of the invention or antigen binding portions thereof, can

antibody, or antibody portion, of the invention can be combined include the following:

corticosteroids; prednisolone; methylprednisolone; azathioprine; cyclophosphamide;

Biogen); interferon-\$1b (Betaseron; Chiron/Berlex); Copolymer 1 (Cop-1; Copaxone;

Teva Pharmaceutical Industries, Inc.); hyperbaric oxygen; intravenous immunoglobulin;

clabribine; antibodies to or antagonists of other human cytokines or growth factors, for

example, TMF, LT, IL-1, IL-2, IL-6, IL-7, IL-8, IL-15, IL-16, IL-18, EMAP-II, GM
example, Topic antibodies of the invention, or antigen binding portions thereof,

can be combined with antibodies to cell surface molecules such as CD2, CD3, CD4,

can be combined with antibodies to cell surface molecules such as CD2, CD3, CD4,

be combined with IL-11.

Non-limiting examples of therapeutic agents for multiple sclerosis with which an

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CD8, CD25, CD26, CD26, CD40, CD45, CD69, CD80, CD86, CD90 or their ligands. The antibodies of the invention, or antigen binding portions thereof, may also be combined with agents, such as methotrexate, cyclosporine, FK506, rapamycin, mycophenolate mofetil, leflunomide, NSAIDs, for example, ibuprofen. corticosteroids auch as prednisolone, phosphodiesterase inhibitors, adensosine agonists, antithrombotic by proinflammatory cytokines such as TMFα or IL-1 (e.g. IRAK, MIK, IKK, p38 or by proinflammatory cytokines such as TMFα or IL-1 (e.g. IRAK, MIK, IKK, p38 or selectin glycoprotein ligand (PSGL), TACE inhibitors, T-cell signalling inhibitors such as kinase inhibitors, metalloproteinase inhibitors, sulfasalaxine, azathioprine, 6-metaplopurines, angiotensin converting enzyme inhibitors, soluble cytokine receptors and derivatives thereof (e.g. soluble p55 or p75 TNF receptors, sIL-1RI, sIL-1RII, sIL-1RIII, sIL-1RII, sIL-1RII, sIL-1RII, sIL-1RII, sIL-1RII, sIL-1RII, sI

Preferred examples of therapeutic agents for multiple sclerosis in which the antibody or antigen binding portion thereof can be combined to include interferon- β , for example, IFN β 1a and IFN β 1b; copaxone, corticosteroids, IL-1 inhibitors, TNF inhibitors, and antibodies to CD40 ligand and CD80.

6R, soluble IL-13 receptor (sIL-13)) and antiinflammatory cytokines (e.g. IL-4, IL-10,

The pharmaceutical compositions of the invention may include a "therapeutically offective amount" of an antibody or antibody portion of the invention. A "therapeutically effective amount" refers to an amount effective, at dosages and for periods of time necessary, to achieve the desired therapeutic according to factors such as the disease state, age, sex, and weight of the individual, and the antibody or antibody portion to elicit a desired response in the individual. A therapeutically effective amount is also one in which any toxic or individual. A therapeutically effective amount is also one in which any toxic or detrimental effects of the antibody or antibody portion are outweighed by the therapeutically beneficial effects. A "prophylactically effective amount" refers to an amount effective, at dosages and for periods of time necessary, to achieve the desired prophylactic result. Typically, since a prophylactic dose is used in subjects prior to or at prophylactic result. Typically, since a prophylactic dose is used in subjects prior to or at

an earlier stage of disease, the prophylactically effective amount will be less than the

therapeutically effective amount.

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IL-13 and TGFb).

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Dosage regimens may be adjusted to provide the optimum desired response (e.g.,

a therapeutic or prophylactic response). For example, a single bolus may be administered, several divided doses may be administered over time or the dose may be administered over time or the dose may be administered over time or the dose may be suministered over time or the dose may be situation. It is especially advantageous to formulate parenteral compositions in dosage unit form for ease of administration and uniformity of dosage. Dosage unit form as used berein refers to physically discrete units suited as unitary dosages. Dosage unit form as used subjects to be treated; each unit containing a predetermined quantity of active compound subjects to be treated; each unit containing a predetermined quantity of active compound dictated by and directly dependent on (a) the unique characteristics of the invention are dictated by and the particular therapeutic or prophylactic effect to be achieved, and (b) compound and the particular therapeutic or prophylactic effect to be achieved, and (b) the limitations inherent in the art of compounding such an active compound for the treatment of sensitivity in individuals.

effective amount of an antibody or antibody portion of the invention is 0.01-20 mg/kg, more preferably 1-10 mg/kg, even more preferablu 0.3-1 mg/kg. It is to be noted that dosage values may vary with the type and severity of the condition to be alleviated. It is to be further understood that for any particular subject, specific dosage regimens should be adjusted over time according to the individual need and the professional judgment of the person administering or supervising the administration of the compositions, and that dosage ranges set forth herein are exemplary only and are not intended to limit the scope or practice of the claimed composition.

An exemplary, non-limiting range for a therapeutically or prophylactically

VII. Uses of the Antibodies of the Invention

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Given their ability to bind to hIL-12, the anti-hIL-12 antibodies, or portions thereof, of the invention can be used to detect hIL-12 (e.g., in a biological sample, such as serum or plasma), using a conventional immunoassay, such as an enzyme linked immunosorbent assays (ELISA), an radioimmunoassay (RIA) or tissue immunosorbent assays (ELISA), an radioimmunoassay (RIA) or tissue biological sample comprising contacting a biological sample with an antibody, or antibody portion, of the invention and detecting either the antibody (or antibody portion)

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bound to hIL-12 or unbound antibody (or antibody portion), to thereby detect hIL-12 in the biological sample. The antibody is directly or indirectly labeled with a detectable substance to facilitate detection of the bound or unbound antibody. Suitable detectable substances include various enzymes, prosthetic groups, fluorescent materials, horseradish peroxidase, alkaline phosphatase, β-galactosidase, or acetylcholinesterase; cxamples of suitable prosthetic group complexes include streptavidin/biotin and avidin/biotin; examples of suitable fluorescent materials include unbelliferone.

Autorescein, fluorescein isothiocyanate, rhodamine, dichlorotriazinylamine fluorescein, fluorescein, fluorescein isothiocyanate, rhodamine, dichlorotriazinylamine fluorescein, dichlorotriazinylamine, dichlorotriaziny

Alternative to labeling the antibody, hIL-12 can be assayed in biological fluids by a competition immunoassay utilizing rhIL-12 standards labeled with a detectable substance and an unlabeled anti-hIL-12 antibody. In this assay, the biological sample, the labeled rhIL-12 standards and the anti-hIL-12 antibody are combined and the amount of labeled rhIL-12 standard bound to the unlabeled antibody is determined. The amount of hIL-12 standard bound to the anti-hIL-12 antibody is determined. The amount of hIL-12 standard bound to the anti-hIL-12 antibody.

The Y61 and J695 antibodies of the invention can also be used to detect IL-12 from primates. For example, Y61 can be used to detect IL-12 in the cynomolgus monkey, thesus monkey, and baboon. However, neither antibody cross reacts with mouse or rat IL-12 (see Example 3, subsection F).

The antibodies and antibody portions of the invention are capable of neutralizing hIL-12 activity in vitro (see Example 3) and in vivo (see Example 4). Accordingly, the antibodies and antibody portions of the invention can be used to inhibit IL-12 activity, e.g., in a cell culture containing hIL-12, in human subjects or in other mammalian subjects having IL-12 with which an antibody of the invention cross-reacts (e.g. primates such as baboon, cynomolgus and rhesus). In a preferred embodiment, the invention provides an isolated human antibody, or antigen-binding portion thereof, that neutralizes the activity of human IL-12, and at least one additional primate IL-12 neutralizes the activity of human IL-12, and at least one additional primate IL-12

selected from the group consisting of baboon IL-12, marmoset IL-12, chimpanzee IL-12, cynomolgus IL-12 and rhesus IL-12, but which does not neutralize the activity of the mouse IL-12. Preferably, the IL-12 is human IL-12. For example, in a cell culture containing, or suspected of containing hIL-12, an antibody or antibody portion of the invention can be added to the culture medium to inhibit hIL-12 activity in the culture.

In another embodiment, the invention provides a method for inhibiting IL-12.

activity in a subject suffering from a disorder in which IL-12 activity is detrimental.

As used herein, the phrase "a disorder in which IL-12 activity is detrimental" is antibodies of the invention (e.g., testing of dosages and time courses of administration). the latter, such animal models may be useful for evaluating the therapeutic efficacy of 57 cross-reacts for veterinary purposes or as an animal model of human disease. Regarding administered to a non-human mammal expressing a IL-12 with which the antibody purposes (discussed further below). Moreover, an antibody of the invention can be antibody of the invention can be administered to a human subject for therapeutic hlL-12 (e.g., by administration of hlL-12 or by expression of an hlL-12 transgene). An 50 cross-reacts. Still further the subject can be a mammal into which has been introduced the subject can be a mammal expressing a IL-12 with which an antibody of the invention Preferably, the IL-12 is human IL-12 and the subject is a human subject. Alternatively, or antibody portion of the invention such that IL-12 activity in the subject is inhibited. from such a disorder, which method comprises administering to the subject an antibody ۶i 832). The invention provides methods for inhibiting IL-12 activity in a subject suffering et al., (1998) Am. J. Path 152:667-672; Parronchi et al (1997) Am. J. Path. 150:823-150:823-832; Monteleone et al., (1997) Gastroenterology. 112:1169-1178, and Berrebi Fais et al. (1994) J. Interferon Res. 14:235-238; Patronchi et al., (1997) Am. J. Path. and Rheumalism. 41: 306-314; Bucht et al., (1996) Clin. Exp. Immunol. 103: 347-367; (Windhagen et al., (1995) J. Exp. Med. 182: 1985-1996; Morita et al. (1998) Arthritis IL-12 has been implicated in the pathophysiology of a wide variety of disorders

intended to include diseases and other disorders in which the presence of IL-12 in a subject suffering from the disorder has been shown to be or is suspected of being either responsible for the pathophysiology of the disorder or a factor that contributes to a worsening of the disorder. Accordingly, a disorder in which IL-12 activity is detrimental is a disorder in which inhibition of IL-12 activity is expected to alleviate the

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invention in the treatment of a few non-limiting specific disorders is discussed further disorders described herein. The use of the antibodies and antibody portions of the thereof, can be used for the manufacture of a medicine for treating the diseases or described herein. In another embodiment, the antibodies or antigen binding portions binding portions thereof, can be used in therapy to treat the diseases or disorders which IL-12 activity is detrimental. In one embodiment, the antibodies or antigen anti-IL-12 antibody as described above. There are numerous examples of disorders in plasma, synovial fluid, etc. of the subject), which can be detected, for example, using an suffering from the disorder (e.g., an increase in the concentration of IL-12 in serum, example, by an increase in the concentration of IL-12 in a biological fluid of a subject symptoms and/or progression of the disorder. Such disorders may be evidenced, for

A. Rheumatoid Arthritis:

useful in the treatment of autoimmune diseases. of the invention also can be administered with one or more additional therapeutic agents of the antibody or antibody portion may be beneficial. An antibody, or antibody portion, portion, is administered systemically, although for certain disorders, local administration spondylitis, osteoarthritis and gouty arthritis. Typically, the antibody, or antibody rheumatoid arthritis, juvenile rheumatoid arthritis, Lyme arthritis, rheumatoid antibodies, and antibody portions of the invention can be used to treat, for example, present in the sublining layer of the rheumatoid arthritis synovium. The human Arthritis and Rheumatism 41: 306-314). IL-12 positive cells have been found to be synovial fluids from patients with rheumatoid arthritis (see e.g., Morita et al., (1998) from rheumatoid arthritis patients and IL-12 has been shown to be present in the such as theumatoid arthritis. Inducible IL-12p40 message has been detected in synovia S١

Interleukin-12 has been implicated in playing a role in inflammatory diseases

severity of disease. Treatment with the anti-IL-12 mAb early after onset of arthritis C17.15) prior to arthritis profoundly supressed the onset, and reduced the incidence and 30 treatment of mice with an anti-IL-12 mAb (rat anti-mouse IL-12 monoclonal antibody, In the collagen induced arthritis (CIA) murine model for rheumatoid arthritis,

reduced severity, but later treatment of the mice with the anti-IL-12 mAb after the onset of disease had minimal effect on disease severity.

B. Crohn's Disease

Crohn's disease. Increased expression of IFN-y and IL-12 occurs in the intestinal mucosa of patients with Crohn's disease (see e.g., Fais et al., (1994) J. Interferon Res. [1997] Gastroenterology 112: 1169-1178; Berrebi et al., (1998) Amer. J. Pathol. 150: 823-832; Monteleone et al., (1997) Gastroenterology 112: 1169-1178; Berrebi et al., (1998) Amer. J. Pathol. 152: of colitis, e.g., TMBS induced colitis IL-2 knockout mice, and recently in IL-10 knock-out mice. Accordingly, the antibodies, and antibody portions, of the invention, can be used in the treatment of inflammatory bowel diseases.

15 C. Multiple Sclerosis

at the peak of paralysis or during the subsequent remission period reduced clinical 12 mAb delayed paralysis and reduced clinical scores. Treatment with anti-IL-12 mAb relapsing-remitting EAE model of multiple sclerosis in mice, pretreatment with anti-ILexperimental allergic encephalomyelitis (EAE) models of multiple sclerosis. In a role of IL-12 in multiple sclerosis has been investigated using mouse and rat activation and disease (Balashov et al., (1997) Proc. Natl. Acad. Sci. 94: 599-603). The APCs, which perpetuated the cycle leading to a chronic state of a Th1-type immune 57 Increased secretion of IFN- γ from the T cells led to increased IL-12 production by basis of progressive multiple selerosis leading to a Th1-type immune response. with multiple sclerosis revealed a self-perpetuating series of immune interactions as the IL-12. Investigations with T-cells and antigen presenting cells (APCs) from patients Chronic progressive patients with multiple sclerosis have elevated circulating levels of Exp. Med. 182: 1985-1996, Drulovic et al., (1997) J. Neurol. Sci. 147: 145-150). demonstrated in lesions of patients with multiple sclerosis (Windhagen et al., (1995). J. sclerosis. Expression of the inducible IL-12 p40 message or IL-12 itself can be Interleukin-12 has been implicated as a key mediator of multiple

scores. Accordingly, the antibodies or antigen binding portions thereof of the invention may serve to alleviate symptoms associated with multiple selerosis in humans.

Interleukin-12 has been implicated as an important mediator of insulin-

D. Insulin-Dependent Diabetes Mellitus

dependent diabetes mellitus (IDDM). IDDM was induced in NOD mice by administration of IL-12, and anti-IL-12 antibodies were protective in an adoptive transfer model of IDDM. Early onset IDDM patients often experience a so-called "honeymoon period" during which some residual islet cells produce insulin and regulate blood glucose levels better than administered insulin. Treatment of these early onset patients with an anti-IL-12 antibody may prevent further destruction of islet cells, thereby maintaining an endogenous source of insulin.

15 E. Psoriasis

are incorporated by reference.

Interleukin-12 has been implicated as a key mediator in psoriasis.

Psoriasis involves acute and chronic skin lesions that are associated with a TH1-type cytokine expression profile. (Hamid et al. (1996) J. Allergy Clin. Immunol. 1:225-231;

Turka et al. (1995) Mol. Med. 1:690-699). IL-12 p35 and p40 mRMAs were detected in diseased human skin samples. Accordingly, the antibodies or antigen binding portions thereof of the invention may serve to alleviate chronic skin disorders such psoriasis.

The present invention is further illustrated by the following examples which

should not be construed as limiting in any way. The contents of all cited references, including literature references, issued patents, and published patent applications, as cited throughout this application are hereby expressly incorporated by reference. It should further be understood that the contents of all the tables attached hereto (see Appendix A)

\$100 \$100 \$100 \$100 \$100 \$100 \$100 \$100	Table 1	
10 germline VH dp-30 DP-30 DP-30 DP-30 DP-31 DP-42 DP-43 DP-45 DP	VН3	30
EVOLUESGGGLUOPGGSLRLSCAASGFTFS EVOLUESGGGLUVPPGGSLRLSCAASGFTFS EVOLUESGGGLUPPGGSLRLSCAASGFTFS OVOLUESGGGUPPGGSLRLSCAASGFTFS OVOLUESGGGPPGGSLRLSCAASGFTFS	Family Germline	
VOLVESGGGLVQPGGSLRLSCAASGFTFS VQLVESGGGLVQPGGSLRLSCAASGFTFS VQLVESGGGVVQPGRSLRLSCAASGFTFS	rmline Amino	52
	Acid	
CORTENANT STANDERS ST	Sequences	50
HURRON PGRGLEHVS WVRON PGRGGLEHVS	Numbering	
CDR H2 RTBNKANSYTTEYANSVKG LIBNKANSYTTEYANSVKG LIBNKANSYTTEYANSVKG LIBNKANSYTTEYANSVKG GISH NGGSIGYADSVKG GISH NGGSIGYADSVKG GISH NGGSIGYADSVKG GISH NGGSIGYADSVKG GISH NGGSIGYADSVKG GISH NGGSITYADSVKG RIKSKTDGGTTDYANPVKG RIKSKTDGGTTYADSVKG GV SHNGSRTHYADSVKG GV SHNGSRTHYADSVKG GVI SGGSGTYYADSVKG GVI SGGSTYYADSVKG AII GTGGGTYYADSVKG AII SGRGGSTYYADSVKG AII SGRGGSTYYADSVKG AII SGTGGGTYYADSVKG AII STDGSNKYYADSVKG VII SYDGSNKYYADSVKG VII SYDGSNKY	ording	\$1
	6	01
SREDSKNSLYLONSSLATEDLAVYCAR SREDSKNTLYLONSSLATEDLAVYCAR SREDSKNTLYLONSSLATEDLAVYCAR SREDSKNTLYLONSSLATEDLAVYCAR SREDSKNTLYLONSSLATEDLAVYCAR SREDSKNTLYLONSSLARAEDTALYYCAR SRDNAKNSLYLONNSLARAEDTAVYYCAR SRDNAKNSLYLONNSLATEDTAVYYCAR SRDNAKNSLYLONNSLATEDTAVYYCAT SRDNAKNSLYLONNSLATEDTAVYYCAT SRDNSKNTLYLONNSLATEDTAVYYCAT SRDNSKNTLYLONNSLATEDTAVYYCAT SRDNAKNSLYLONNSLATEDTAVYYCAT SRDNAKNSLYLONNSLATEDTAVYYCAT SRDNAKNSLYLONNSLATEDTAVYYCAT SRDNAKNSLYLONNSLATEDTAVYYCAT SRDNAKNSLYLONNSLATEDTAVYYCAT SRDNAKNSLYLONNSLATEDTAVYYCAT SRDNAKNSLYLONNSLAREDTAVYYCAR SRDNSKNTLYLONNSLAREDTAVYYCAR	VH included for comparison	ς

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SEQ 10 843 843 853 853 853 853 853 853 853 853 853 85	
9¢ rml ine VH B1 B13 B16 B29¢ B20¢ B	
	0€
· · · · · · · · · · · · · · · · · · ·	57
1 CC C C C C C C C C C C C C C C C C C	
VH3 Family Germline Numbering according (Joe9 VH included f. WYRQAPGKGLENVA VI. SY WYRQAPGKGLENVS SI. SS WYRQAPGKGLENVS RI. NS	50
TO Amino Acid for Compariso rogswartyadsvartogswartyadsvarty	SI
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VPPENDIX A

	500 10	2.5
90ne. 110 110 110 110		30
DP1.5 DP1.9 DP1.9 DP1.9 DP1.9 DP1.9		
DPL5 QSVLTQPPSV3AAPGQKVTISC SGSSSNIGNYA.VS W DPL4 QSVLTQPPSV3AAPGQKVTISC SGSSSNIGNYA.VS W DPL4 QSVLTQPPSV3AAPGQKVTISC SGSSSNIGSNT.VN W DPL3 QSVLTQPPSASGTPGQRVTISC SGSSSNIGSNT.VN W DPL3 QSVLTQPPSASGTPGQRVTISC SGSSSNIGSNY.VY W DPL1 QSVLTQPPSVSEAPRQRVTISC TGSSSNIGAGYVVH W DPL9 QSVLTQPPSVSGAPGQRVTISC TGSSSNIGAGYVVH W DPL8 QSVLTQPPSVSGAPGQRVTISC TGSSSNIGAGYVH W DPL8 QSVLTQPPSVSGAPGQRVTISC TGSSSNIGAGYPH W DPL8 QSVLTQPPSVSGAPGQRVTISC TGSSSNIGAGAT QSVLTQPPSVSGAPGAT QSVLTQPSVLTQPT QSVLTQPT QSVLTQPT QSVLTQPT QSVLTQPT QSVLTQPT QSVLTQPT QSVLTQPT QSVLTQPT QSVLTQPT QS		
SGSSSNIGNY. VS SGSSSNIGSNT. VN SGSSSNIGSNT. VN SGSSSNIGSNY. VY SGSSSNIGN. AVN TGSSSNIGAGYVVH TGSSSNIGAGYVVH SGCRSNIGAGYDVH SGCRSNIGAGYDVH	Table 1 V	50
YUKUTOTAPKILIY DNNKRPS YQQIPGTAPKILIY ENNKRPS YQQIPGTAPKILIY SNNQRPS YQQIPGTAPKILIY RNNQRPS YQQIPGTAPKILIY RNNQRPS YQQIPGTAPKILIY GNSNRPS YQQIPGTAPKILIY GNSNRPS YQQIPGTAPKILIY GNSNRPS YQQIPGTAPKILIY GNSNRPS	VAl Family Germline Amino Acid Se Numbering according to Kabat. (Joe9 VL included for comparison)	12
GIPDRESGSKSGTSATLGITGLOTGDEADYYC GTWDSSLSA GIPDRESGSKSGTSATLGITGLOTGDEADYYC GTWDSSLSA GIPDRESGSKSGTSATLGITGLWPEDEADYYC LAWDTSPRA GVPDRESGSKSGTSASLAISGLQSEDEADYYC AAWDDSLNG GVPDRESGSKSGTSASLAISGLRSEDEADYYC AAWDDSLNG GVPDRESGSKSGTSASLAITGLQSEDEADYYC AAWDDSLNG GVPDQESGSKSGTSASLAITGLQSEDEADYYC AAWDDSLNG GVPDRESGSKSGTSASLAITGLQSEDEADYYC QSYDSSLNA GVPDRESGSKSGTSASLAITGLQAEDEADYYC QSYDSSLNA GVPDRESGSKSGTSASLAITGLQAEDEADYYC QSYDSSLNA	no Acid Sequences (abat	01
TC QSYDSELRG	CDR LJ	ς

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		1.00E-09	8.90E-04	QSYDNAVTASKV	123	HGSHDN	83	136-24
		2.00E-09	4.20E-04	QSYDNGFTGSRV	122	HGSHDN	83	136-21
	100	8.00E-10	1.10E-03	QSYDRGFTGSRV	121	HGSHDN	83	136-18
	7.005-09	2.00E-09	2.90E-05	QSYDWNFTGSRV	120	HGSHDN	83	136-17
	5.008-09	3.00E-10	6.10E-04	QSYDRRFTGSRV	119	HGSHDN	83	136-16
	6.005-09		4.60E-04	QTYDKGFTGSSV	118	HGSHDN	83	136-15 germline
	2.00F-09	1.00E-10	7.4 e-4	QTYDKGFTGSSV	118	HGSHDN	83	136-15
	1.00F-07	3.00E-10	1.10E-03	QTYDRGFTGSRV	117	HGSHDN	8.3	136-14
		2.00E-09	1.40E-03	QSYDRGETGSRV	116	HGSHDD	82	136-10
			3.20E-03	QTYDISESGSRV	115	HGSHDD	82	136-9
	6.005-08		7.4 e-3	79-1	114	70-1	78	26-1 (2,3)
		1.60E-09		78-34	111	70-1	78	101-11 IgG1
	3.00E-08		4.5 e-3	78-34	111	70-1	78	101-11 (12)
			4.45E-02	78-35	113	70-13	81	101-5
			9.76E-03	78-35	113	70-13	81	101-4
			1.01E-02	78-34	111	70-13	81	101-8
			4.56E-02	78-34	111	70-13	81	101-19
			8.54E-03	78-35	113	70-2	79	101-9
			7.52E-03	78-34	111	70-2	79	101-14
	6.005-07	2.00E-07		QSYDSSLWGSRV	114	₩t	77	. 79-1
		4.00E-07	4.99E-02	QSYDSSLTGSRV	113	₩t	77	78-35
			4.66E-02	QSYDSSLRGSRV	112	wt	77	78-28
			5.00E-02	QSYDSSLRGSRV	112	₩t	77	78-25
	6.005-07	2.002-07	1.64 e-2	QSYDRGFTGSRV	111	Μť	77	78-34
	3-5 08-7		7.20E-02	Joe9 wt	110	SGSIDY	81	70-13
	3-5 06-7		1.29E-01	Joe9 Ht	110	RRRSNY	80	70-7
	1-505-07		3.30E-02	J089 Wt	110	HGSYDY	79	70-2
	2 005-07			Joe9 wt	110	HGSHDN	78	70-1 IgG1
	2005-07		1.34 e-2	јое9 мt	110	HGSHDN	78	70-1
	5.008-07			QSYDSSLRGSRV	110	SGSYDY	. 77	Joe9 wt IgGl
1C20 (H)	1 005-06	1.50E-06	1.00E-01	VASSUASSOASO	110	SGSYDY	77	Joe9 wt
	ICSO (M)	IC50 (M)	koff	1.3	ID NO:	нз		Clone
7	PHA	RB assav			L3 SEQ		H3 SEQ	
				Table 2				
Ş		01		ŞI	50		57	09

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156-18	156-17	156-16	156-15	156-14	156-13	156-12	156-11	156-10	156-9	156-8	156-7	156-6	156-5	156-4	156-3	156-2	156-1		149-14	149-13	149-12	149-11	149-9	149-8	149-7	149-6	149-5	149-4	136-15M1	101-11	Clone		
93	92	93	94	94	94	94	94	92	92	92	92	92	93	93	93	93	93	92	91	84	8.4	90	89	88	84	84	87	86	85		H3 SEQ		
			. X	. 🛪	. ×	. 🛪	. K								T		T	TT HGSHDN	RN			S	.				. 1	.		IT HESHIN HEOG	нз		
125	125	124	130	12/	128	521	120	130	12,	128	129	126	130	127	128	129	126	124				128	127		126	124	125	124	124	124	L3 SEQ		,
SSLW.1	SSLW.1		.T		:		١:	.1		1:			.TK	1:	2	1 .	D	OSYDRGETGSRY				NA	E		D		QSYDSSLWGTRV		QSYDRGFTGSRV	OSYDRGETGSRY	Г.3	Table 2	
	6.00E-03	202				+						3.006-03			9.006-03		5.00E-03			2.22×10-3		1.43×10-2	3.54×10-3	2.33×10-3	1.13×10-3	73×10-	1.02×10-3	1.37×10-3		4.5×10-3	koff		
																			1.5×10-10		1_	\vdash	1-	1		6×10-10	1.2×10-10	8×10-11	4.00E-10	2X10-9	RB assay IC50 (M)		C
																			6.00E-09		ing	4.00E-09			3.00E-09	2.00E-09	3.00E-09	3.00E-09		2.00E-08	PHA assay ICSO (M)		
																															IFN gamma		ç

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П																					-		1					İ						
	170-25	170-36	170-39	170-38	170-35	170-24	170-23	170-22	170-21	170-19	170-15	170-13	170-11	170-7	170-4	170-3	170-2	170-1		103-152	103-4	103-8 & 2	103-14 & 9	103-8	103-7	103-6	103-3	103-2	103-1		Clone			30
	106	83			105	104	103	102	104	103	102	102	102	102	102	102	102	102	102	101	101	100	100	99	98		97	96	95	92	H3 SEQ			
	HGSQDT	HGSHDN			A. HQ.N	H. Q.N	HH.N		HQ.N	HH.N									TI SGSXDX	TT HGSHDN	TT HGSHDN	KT HGSHDN	KT HGSHDN	K	D		· · · · · · · · · · · · · · · · · · ·	K. R.R	Q.R	TT HGSHDN	нз			\$7
-	153	152	151	150	136	149	148	147	146	145	144	143	142	141	140	139	138	137	136	135	134	133	132	130	131	131	124	130	124	124	L3 SEQ ID NO:			07
	CSTDSSTROSKAL	QSYDRUSTGSKVE	м. S.	P		Y					LY.L.	YK	L F		Y.A	L. VIX	VSAY		OSYDRGETGSRYE	QSYERGETGARV	QSYDRGFTGARV	QTYDKGFTGSSV	QSYDRGFTGSMV	.TK S.	DT	DT		.TKS.		OSYDRGETGSRY	1.3	Table 2	1	ŞI
	3.00E-04	4.00E-04	2.79E-03	2.10E-04		2.80E-04	١.	5.60E-04	3.89E-03	1.005-03	4.43E-03	1.59E-03	4.40E-04	١.	1.	1.		2.35E-03			1.6 e-4	i.u	1~	3.3×10-4		4.5×10-4	2.5×10-3	7.3×10-4	2.9×10-3		koff			
	5.00E-11	2.00E-10				5.00E-10	1.													8.60E-11	8.60E-11	1	4.00E-11	6.00E-11	1.40E-10			7.00E-11			RB assay IC50 (M)			01
																					9.00E-10	1.50E-09	1.20E-09	1.50E-09	1.00E-09			1.00E-09			PHA assay IC50 (M)			
																															IFN gamma			ç

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73-B7	73-C5	73-Н3	73-86	73-н2	73-G6	73-65	73-64	73-G3	73-G2	73-F5	73-F3	73-E6	73-E3	73-DS	73-D4	73-D2	73-D1	73-C6	73-C2	73-C1	73-B6	73-B2	73-B1		Clone					30
108	107	107	107	107	107	107	107	107	107	107	107	107	107	107	107	107	107	107	107	107	107	107	107	106	H3 SEQ					52
HGSQDN	SGSYDY	SGSYDY	SGSYDY	SGSYDY	SGSYDY	SGSYDY	SGSYDY	SGSYDY	SGSYDY	SGSYDY	SGSYDY	SGSYDY	SGSYDY	SGSYDY	SGSYDY	SGSYDY	SGSYDY	SGSYDY	SGSYDY	SGSYDY	SGSYDY	SGSYDY	SGSYDY	SGSYDY	нз					
177	176	175	174	173	172	171	170	169	168	167	166	165	164	163	162	161	160	159	158	157	156	155	154	136	L3 SEQ					50
QSYDSSLRGSRV	н.н D	HN	HM.	S	NTD	H.S.SDS	MR	HTR	HTN.S	DSDK	APWS	HES	SD	H.S.S	HTK	T	HS	T	SE	HSR	HNR	H.SES	HSD	OSYDRGETGSRYE	1.3	Table 2	1			ŞI
2.50E-03	4.84E-03	6.85E-03	5.87E-03	5.93E-03		6.01E-03		3.50E-03	3.98E-03	3.74E-03	7.08E-03	4.17E-03	4.98E-03	3.57E-03	5.36E-03	3.56E-03	3.99E-03	3.96E-03	3.79E-03	2.71E-03	2.51E-03	2.07E-03	3.25E-03		koff					
7.00E-09																				>1E-8	>1E-8		>1E-8		RB assay IC50 (M)					01
																									PHA assay				,	
																								(H) 0531	IEN gamma					Š

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			9.90E-03	A	198	HGSHDN	83	м2 н5	
			6.16E-02	SDV	197	HGSHDN	83	M2 F1	
			9.77E-03	TX	196	HGSHDN	83	M2 E6	
			4.77E-02		195	HGSHDN	83	M2 E2	
			5.32E-02	.T. X S.	194	HGSHDN	83	M2 E1	
			5.01E-02	SA	193	HGSHDN	83	1	
			5.18E-02	SSS	192	HGSHDN	83	H2 D5	
			5.81E-02	ETS	191	HGSHDN	83	M2 D4	
			3.33E-02	NRL	190	HGSHDN	83	M2 D3	
			3.85E-02	IRS	189	HGSHDN	83	H2 D2	
			3.71E-02	IR	188	HGSHDN	83	M2 D1	
			4.71E-02	TAL	187	HGSHDN	8.3	M2 C5	
			8.16E-03	S.L	181	HGSHDN	83	- 1	
				T.L	186	HGSHDN	83	H2 C3	
			4.85E-02	S.L	181	HGSHDN	83	M2 C2	
			2.81E-02	L.A	185	HGSHDN	83		
			8.38E-03	S.V	184	HGSHDN	83	M2 B5	
			4.42E-02	I.L	183	HGSHDN	83	M2 B4	
			4.60E-02	I.M	182	HGSHDN	83	M2 B3	
		•	7.97E-03	S.L	181	HGSHDN	83	M2 B1	
			4.01E-02	I.S	180	HGSHDN	83	M2 A5	
			8.49E-03	SP	179	HGSHDN	83	M2 A4	
			4.00E-02	IH	178	HGSHDN	83	M2 A2	
7				OSYDRGETGSRVE	136				
ICSO (M)	IC50 (M)	IC50 (M)	koff	13		н3		Clone	
	PHA assay	RB assay			L3 SEQ		H3 SEQ		
				Table 2					
~	ζ.								
Ş		0 (51	07	57		30	
		1							

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			89 8	A9 8	00			C12		AI		B7		В3		A7		A2		C3				BS		A3	9.0	88	A8	C2	All	A10	A6	A4	AIZ	AS		_				
			٦	ω	- ω					83				~ "	+	2	1	83	83	1		_	-	83			B) (R. C		8	83	83	83	83	83	83		NO:	H3 SEQ			
		TO TO THE	COHON	HGSHDN	HGSHDN	HGSHDN	MONHON	00000	HOSHDN	HGSHDN	HGSHDN	HGSHDN	HOUHUN	TOUT ON	NOHON	NOUNDE	DOCUMENT OF THE PARTY OF THE PA	HCGHON	HOSHON	HGSHDN	HGSHDN	HGSHDN	HGSHDN	HGSHDN	HGSHDN	HGSHDN	NOHON	HOGH DN	HOSHON	accino.	HOSHON	HGSHDN	HGSHDN	HGSHDN	HGSHDN	HGSHDN		нз				
		. 677	<u> </u> -	278	227	226	225	. 677	3 :	222	222	221	220	612	817	217	917	21.0	315	214	212	212	211	210	209	208	207	206	205	204	203	202	303	315	300	198	124		L3 SEQ			
		RAAHPQ	PARTSP	1100000	119915	4	LRVQAP	TYSTAI	H.T. TAKT. H	TOWNED L	משמחו	S. TSNLIP	TPSYPT	HATHTA	HNNFSP	PHTQPT	PYHPIR	RPRHAL	YAPANA . C	١.	171 200	Taldon	OPHAVI	ITPGLA	YPRNIL	QTPSIT	HTAHLY	KSNKML	LNPSAT	TFPSPQ	NSPATV	HIMAHI	KNPALT	ITERPM	TWOME	AMERICAN SECTIONS	DOWNER	1.3		Table 2	,	
		8.66E-03	7.98E-03	7.588-03	0.048-03		5.85E-03	5.83E-03	5.66E-03	5.45E-03	20.000.03	2000	4.12E-03	3.97E-03	3.95E-03	3.91E-03	3.80E-03	3.61B-03	3.34E-03	1 .	3. TOE-03	101E-03	3 0 0 0 0 0		2 51E-03	2.42E-03	2.40E-03	2.37E-03	2.23E-03	2.07E-03	1.876-03	1.87E-03	1.478-03	1.438-03	┼	1	YOLL					
	1	1	1		<u> </u>			-	1		-							$\frac{1}{1}$				 						-				\downarrow	$\frac{1}{1}$				_	ssay				
					$\left \cdot \right $		 -			$\left \right $		\int		 -	 																					(1)	ICSO (M)	PHA assay				
					. -																 								\int							ICSO (M)	LEN gamma					

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	>1E-8	1.40E-10	1.006-03	Car Divorus Aut	737	HGSHUN	90	133
	1.00E-08	3.5E-10	1.07E-03	QSYDRGQTPSIT	256	HGSHDN	83	C6
	1.005-09			THPLTI	255	HGSHDN	83	G6 IgG1
 	1.00E-09	1E-10	5.08E-04	THPLTI	255	HGSHDN	83	66
	×	3.5E-09	8.21E-04	RLPAQT	254	HGSHDN	83	D9
		2.50E-10	5.57E-04	TTPRPM	253	HGSHDN	83	A12
	- 100	6.60E-11	4.50E-04	THPSML	252	HGSHDN	83	A5
	1.005-00	5E-11	6.61E-04	SHPTLI	251	HGSHDN	83	7177
	2.005-09	6E-11	7.55E-04	TTPAPE	250	HGSHDN	83	Y174
	1 005-00			SHPALT	249	HGSHDN	83	Y139 IgG1
1.60E-10	4.508-10	3E-11	5.92E-04	SHPALT	249	HGSHDN	83	Y139
	1.30F-10	1.60E-11	1.50E-04	THPALL	248	HGSHDN	83	Y61 IgG germline
	1 305 10	1.60E-11	1.50E-04	THPALL	248	HGSHDN	83	Y61 19G
	405-10	4E-11	2.75 e-4	THPALL	248	HGSHDN	83	Y61
	4 305-00	4.00E-11	6.17E-04	TDPAIV	247	HGSHDN	83	Y 4 5
	2 605-00	5.50E-11	5.08E-04	THPSIT	246	HGSHDN	83	¥38
	9-310	3.00E-11	2.26E-04	QSYDRGHHPAMS	245	HGSHDN	83	Y19
	>15-8	4.50E-10	8.90E-05	QSYDRGSAPMIN	244	HGSHDN	83	Y17
				OSYDRGETGSRY	124			
			5.00E-04	THPSIS	243	HGSHDN	8.3	31-B5
			5.00E-04	THPNLN	242	HGSHDN	8.3	31-A6
			9.00E-04	IWPNLN	241	HGSHDN	83	43-G3
			7.00E-04	SSPAIM	240	HGSHDN	83	43-G2
			8.00E-04	TIPSIE	239	HGSHDN	83	43-E9
		•	8.00E-04	SHPAAE	238	HGSHDN	8.3	43-E3
			5.80E-04	HHYTTE	237	HGSHDN	83	144-F1
			3.44E-03	THPTMY	236	HGSHDN	6.8	177-H10
-			1.58E-03	THPVPA	235	HGSHDN	£8	177-H9
			1.32E-03	SBPIPA	234	HGSHDN	6.8	177-H5
			7.94E-04	THPVPA	233	HGSHDN	83	177-C6
			6.32E-04	TEAGIA	232	HGSHDN	83	177-D9
			5.50E-04	THPTMI	231	HGSHDN	83	177-66
			4.07E-04	TQPABI	230	HGSHDN	83	177-D7
IC50 (M)	(B)			OBYDRGETGSRY	124			
IFN gamma	PHA assay	RB assay	koff	Г3	ID NO:	н3	H3 SEQ	Clone
				Table 7				
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			3.95E+00	MIPALT	282	HGSHDN	83	. 1
			4.24E+00	FTGSMV	281	HGSHDN	8.3	L3.3R3M-G7
			4.21E+00	FTGSMV	280	HGSHDN	8.3	. 1
			4.99E+00	TTPRIR	279	HGSHDN	83	.
			5.37E+00	FTGSMV	278	HGSHDN	8.3	L3.3R3M-A3
			5.35E+00	TAPSLL	277	HGSHDN	83	L3.3R3M-H10
			5.69E+00	NWPNSN	276	HGSHDN	83	L3.3R3M-H6
			5.55E+00	SYPALR	275	HGSHDN	83	
			4.99E+00	TAPALS	274	HGSHDN	83	L3.3R3M-F9
			6.17E+00	FTGFDG	273	HGSHDN	83	
			5.51E+00	FTGSMV	272	HGSHDN	8.3	L3.3R3M-B3
			٠.	FTGSMV	271	HGSHDN	83	L3.3R3M-B1
				OSYDRGETGSRY	124			
			4.30E-03	QSYDSRFTGSRV	270	HGSHDN	6.8	99-G11
			4.80E-03	QSYLKSRAFSRV	269	HGSHDN	8.3	99-G7
			5.40E-03	QSYDSGFTGSRV	262	HGSHDN	83	99-F11
			3.70E-03	QSYIRAPQQV	268	HGSHDN	8.3	99-F8
			4.80E-03	QSYPNSYPISRV	267	HGSHDN	83	99-F7
			4.90E-03	QSYDSGSTGSRV	266	HGSHDN	83	93-66
			4.90E-03	QSYSTHMPISRV	265	HGSHDN	83	99-н11
			3.30E-03.	QSYPDGTPASRV	264	HGSHDN	83	99-H7
			5.40E-03	QSYDSGFTGSRV	262	HGSHDN	83	99-E9
			4.80E-03	QSYDSRFTGSRV	263	HGSHDN	8.3	99-H4
			5.70E-03	QSYDSGETGSRV	262	HGSHDN	83	99-C11
			5.40E-03	QSYDSGYTGSRV	261	HGSHDN	83	99-811
	100							
	755-10	.50E-	2.50E-04	QSYDRGTHPLTM	260	HGSHDN	83	
	3 808-10	2.90E-11	3.04 e-4	MITAHLDAGASÖ	260	HGSHDN	83	AO3 IgG1
	4.00E-10	3.00E-11	3.04E-04	QSYDRGTHPLTM	259	HGSHDN	83	A03
1C30 (M)	- 1	2.50E-10	6.30E-04	QSYDRGRNPALT	258	HGSHDN	83	A4
IFN gamma	PHA assay	RB assay IC50 (M)	koff	£.1	ID NO:	H3	H3 SEQ	Clone
				Table 2				
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j		to E	н1: н318	*CDR L2: L50G to Y; CDR H1: H31S to E	*CDR L2: L50G to Y	
5.00E-11	5.14E-05	QSYDRYTHPALL	287	CKT HGSHDN	109	L50Y IgG*)
	4.64E-05	QSYDRYTHPALL	287	CKT HGSHDN	109	194Y** IgG
	2.99E-05	QSYDRGTHPALL	286	CKT HGSHDN	109	IgG IgG
+	6.98E-05	QSYDRGTHPALL	286	CKT HGSHDN	109	161-150Y* IgG
+		QSYDRGTHPALL	286	CKT HGSHDN	109	Y61-L50Y
	1.27E-04	QSYDRYTHPALL	285	CKT HGSHDN	109	Y61-L94Y IgG
+		QSYDRYTHFALL	285	CKT HGSHDN	109	Y61-L94Y
		QSYDRFTHPALL	284	CKT HGSHDN	109	761-L94F
+	7011	QSYDRNTHPALL	28-3	CKT HGSHDN	109	Y61-L94N
	K 0 1	1.3	ID NO:	н 3	ID NO:	Clone
1 1		Table 2				

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tion	contact and/or hypermutation position	or hyper	x contact and/o	
×	×	××	×	Hypermutation Positions
×	×××	XXXX	×	Contact Positions
D 0	GNDQRPS	N T V K	SGGRSNIGS	Y61 VL
68 06 16 26 6 6 6	0S TS 2S ES \$S \$S	35 35 36 36	Kabat number	Kabat number
CDR L3	CON LZ		2 2	
	2 400		CDR 1.1	
	X X	×	XXX	Hypermutation Positions
	X	XXX	X X X X	Contact Positions
A D S V K	DGSNKYY	FIRY	FTFSSYGMH	Y61 VH F T
09 19 29 29 9 9 6 6	ES FS SS 9S LS 8S 6S	ASS SS IS 08	Kabat Number 2226669 SEE SEE SEE SEE SEE SEE SEE SEE SEE SE	Kabat Number
	CDR H2		CDR H1	
		Table 3	Tab	

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Table 4	Neutralizati	on Activity in the	Neutralization Activity in the Presence of Excess Free IL-12	Free IL-12 p40
SEQ ID NO:	Clone	PHA assay IC50 (M) p70:p40 1:0	PHA assay IC50 (M) p70:p40 1:20	PHA assay IC50 (M) P70:p40 1:50
VH: 47	136-15	2.00E-09	5.00E-09	
VL: 48				
VH: 51	149-5	6.50E-09	7.00E-09	4.00E~00
VL: 52	,			
VII: 53	149-6	9.00E-10	1.00E-09	l . 00E - 09
VL: 54				
VH: 84	149-7	3.50E-09	2.50E-09	4 007-00
VL: 126				. 005-09
VH: 23	761 IgG	1.808-10		
VL: 24				1.80E-10
VH: 65	A03 [qG]	2.50E-10		2.205-10
VL: 66				
VH: 31	J695	1.00E-11		3.50E-11
VL: 32				

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EXYMPLES

Isolation of Anti-IL-12 Antibodies EXYMDEE 1:

A. Screening for IL-12 binding antibodies

(PBL) (referred to as seFv 2), and bone marrow-derived lymphocytes (referred to as from human tonsils (referred to as scFv I), tonsil and peripheral blood lymphocytes phage display libraries prepared using human VL and VH cDNAs from mRNA derived Antibodies to hIL-12 were isolated by screening three separate scFv

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et al. (1996) Nature Biotech. 14: 309-314. BMDL). Construction of the library and methods for selection are described in Vaughan

by coating the antigen onto immunotubes using standard procedures (Marks et al., human IL-12 and biotinylated chimaeric IL-12. IL-12 specific antibodies were selected IL-12 p40 subunit, chimaeric IL-12 (mouse p40/human p35), mouse IL-12, biotinylated The libraries were screened using the antigens, human IL-12 p70 subunit, human

- VHDP58/VLDPL11, VHDP77/VLDPK31, VHDP47/VL and VHDP77/VLDPK31, all of patterns, and confirmed by DNA sequencing. The main clonotypes were Five different clonotypes were selected, determined by BstMI enzymatic digestion or biotinylated-IL-12, and generated a significant number of IL-12 specific binders. (1991) J. Mol. Biol. 222: 581-597). The scFv library 2 was screened using either IL-12,
- Screening of the BMDL library with IL-12 p70 generated 3 different clonotypes. which recognized the p40 subunit of IL-12.

IL-12. Screening of the scFv library 1, using IL-12 p70, did not produce specific IL-12 sequenced and consisted of VHDP35/VLDP. This clone recognizes the p40 subunit of Two of these were found to be cross-reactive clones. The dominant clone was

antibody lineages. Subunit preferences were further analyzed by 'micro-Friguet' selected in the presence of free p40. Sequencing of isolated clones revealed 9 different recognized the p70 heterodimer or p35 subunit, phage libraries were preincubated and 1 + 2 library, and the BMDL library were used. To select IL-12 antibodies that heterodimer or the p35 subunit of IL-12, rather than the p40 subunit, the combined scFv In order to identify IL-12 antibodies which preferentially bind to the p70 antibodies.

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and p70. Any variation from this gives the degree of preference of p70 over p40. p70 and p40. If the titrations for both subunits overlaps, then the scFv binds to both p40 concentration of free p70 or p40. This provided the IC50 for each clone with respect to in the ELISA signal on biotin-IL-12 coated plates was measured and plotted against the preincubated with increasing concentrations of free p70 or p40 (inhibitors). A decrease ELISA and the ED50 determined. The concentration of scFv producing 50% ED was titration. The supernatant containing scFv was titrated on biotin-captured IL-12 in an

affinity maturation. The amino acids sequences of VH and VL of the Joe9 wild type members of $V_{k}I$ family of germline sequences. Therefore, loe 9 was selected for shows the V_H3 family of germline sequences, of which COS-3 is a member, as well as sequence COS-3, identified from the VBASE database. Table I (see Appendix A) engion (VH) of Joe 9 had the least number of changes compared to the closest germline with an IC₅₀ value of 1 x 10.0 M in both assays. In addition the heavy chain variable Example 3. Clone Joe 9 had the lowest IC_{50} value in both the RBA and the PHA assay, 12 induced proliferation of PHA stimulated human blast cells (PHA assay), described in an IL-12 receptor binding assay (referred to as RBA), and for their ability to inhibit IL-01 The clones were tested for their ability to inhibit IL-12 binding to its receptor in B.

Affinity Maturation of Antibody Lineage Specific for IL-12 (Joe 9)

containing a random mixture of all four nucleotides, chain CDR3 was performed using the degenerate heavy chain oligonucleotide substitutions in each CDR3 (referred to as "spike"). PCR mutagenesis of the heavy "H3") or the light chain CDR3 (referred to as "L3"), with an average of three base degenerate oligonucleotides specific for either the heavy chain CDR3 (referred to as made. The CDR3 variants were created by site-directed PCR mutagenesis using complementarity determining region 3 (CDR3) of both the heavy and light chains were In order to increase the affinity of Joe 9, various mutations of the

(loe9 wt) antibody are shown in Figure 1A-1D.

GAC ACC TCG ATC AGC GGA TAA CAA TTTCAC ACA GG (SEQ ID NO: 581) GGTCGTACAGTAATA 3' (SEQ ID NO: 580), and oligonucleotide pUC Reverse Tag

to generate a repertoire of heavy chain CDR3 mutants. The parent light chain was amplified using Joe 9 reverse oligonucleotide (5'TGG GGC CAA GGG ACA3' (SEQ ID NO:582) and the fdteteseq 24+21 oligonucleotide (5'-ATT CGT CCT ATA CCG TTC TAC TTT GTC GTC TTT CCA GAC GTT AGT-3' (SEQ ID NO: 583).

Complementarity between the two PCR products was used to drive annealing of

the two fragments in a PCR assembly reaction and the full length recombined scFv library was amplified with pUC Reverse Tag (SEQ ID NO: 584) and fdTag 5'-ATT CGT CCT ATA CCG TTC-3' (SEQ ID NO: 584). PCR mutagenesis of the light chain was performed using the light chain oligonucleotide containing a mixture of all four nucleotides

5'GGTCCCAGTTCCGAAGACCCTCGAACC(C)(C)(T)(C)(A)(G)(G)(G)(C)(T)

(G)(C)(T)(G)(T)(C)ATATGACTGGCAGTAATAGTCAGC 3' (SEQ ID NO: 585), and loe 9 reverse oligonucleotide 5'TGG GGC CAA GGG ACA3' (SEQ ID NO: 586)

amplified with pUC Reverse Tag (SEQ ID NO: 581) and HulH3FOR oligonucleotide 5'TGAAGACGTGACCATTGTCCC3' (SEQ ID NO: 587). Complementarity between the two PCR products was used to drive annealing of the two fragments in a pctween the two PCR products was used to drive annealing of the two fragments in a pctween the two PCR products as used to drive annealing of the two fragments in a hotween the two PCR products as used to drive annealing of the two fragments in a pctween the two PCR products as used to drive annealing of the two fragments in a hot well as two PCR products as used to drive annealing of the two fragments in a pctween the two PCR products as used to drive annealing of the two fragments in a pctween the two PCR products as used to drive annealing of the two fragments in a pctween the two PCR products as used to drive annealing of the two fragments in a pctween the two PCR products as used to drive annealing of the two fragments in a pctween the two PCR products as used to drive annealing of the two fragments in a pctween the two PCR products as used to drive annealing of the two fragments in a pctween the two PCR products are used to drive annealing of the two fragments and the following products are used to drive annealing of the two fragments and the following products and the following products are used to drive annealing products and products and products are used to drive annealing products and products and products and products are used to drive annealing products and products are used to drive annealing products and products are used to drive annealing products and products are used to drive annealing products and products are used to drive annealing products and products are used to drive annealing products and products are used to drive annealing products and products are used to drive anneali

GGT CAG CTT GGT CCC 3' (SEQ ID NO: 589).

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Heavy chain CDR3 mutants were selected using 1 nM biotinylated IL-12, and washed for 1 h at room temperature in PBS containing free IL-12 or p40 at a concentration of 7 nM. Clones were analyzed by phage ELISA and those that bound to procedure for BIAcore kinetic binding studies using a low density IL-12 chip (see real-time binding interactions between ligand (recombinant human IL-12 immobilized on a biosensor matrix) and analyte (antibodies in solution) by surface plasmon resonance on a biosensor matrix) and analyte (antibodies in solution) by surface plasmon resonance utilizes the optical properties of SPR to detect alterations in protein concentrations within a dextran biosensor matrix. Proteins are covalently bound to the dextran matrix at known concentrations. Antibodies are injected through the dextran matrix and at known concentrations. Antibodies are injected through the dextran matrix and

specific binding between injected antibodies and immobilized ligand results in an increased matrix protein concentration and resultant change in the SPR signal. These changes in SPR signal are recorded as resonance units (RU) and are displayed with respect to time along the y-axis of a sensorgram. To determine the off rate (k_{off}), on rate (k_{on}), association rate (Ka) and dissociation rate (Kd) constants, BIAcore kinetic evaluation software (version 2.1) was used. Clones that demonstrated an improvement in the k_{off} rate were analyzed by neutralization assays which included inhibition by proliferation in PHA stimulated human blast cells (PHA assay), and inhibition of IL-12-induced induced interferon gamma production by human blast cells (IFM gamma assay). A induced interferon gamma production by human blast cells (IFM gamma assay). A

summary of the dissociation rates and/or IC $_{50}$ values from neutralization assays of heavy chain CDR3 spiked clones 70-1 through 70-13 is presented in Table 2 (see Appendix A). Clone 70-1 displayed a $k_{\rm off}$ rate that was better than the parent Joe 9 clone, and had the lowest IC $_{50}$ value of 2.0 x 10^{-7} M. Therefore clone 70-1 was selected for conversion to complete IgG1.

Light chain CDR3 mutants were selected using 1 nM biotin-IL-12 and washed with PBS containing 7 nM free p40. Clones were screened in phage ELISA and those that bound to IL-12 were tested in BIAcore binding analysis using low density IL-12 were tested in neutralization assays which measured either, inhibition of IL-12 receptor binding, or inhibition of PHA blast cell proliferation. A summary of the dissociation rates and/or IC₅₀ values from neutralization assays of light chain CDR3 mutant clones, 78-34 through 79-1, is presented in Table 2 (see Appendix A).

Based on the k_{off} rate, clones 78-34 and 78-35 displayed an improved k_{off} rate

compared to the parent Joe 9. Both of these clones were selected for combination analysis with heavy chain mutants.

Combination Clones

Mutant light and heavy chain clones that exhibited the best binding

ocharacteristics were used for combination and assembly of scFvs. Mutant clones with

improved potency characteristics were combined by PCR overlap extension and pull-

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through 26-1, shown in Table 2 (see Appendix A), were produced from the combination of heavy chain mutants (70-2, 70-13 and 70-1) with light chain mutants (78-34, 78-35 and 79-1). The k_{off} rates and/or IC₅₀ values from neutralization assays for these clones are presented in Table 2.

BIAcore binding analysis identified clone 101-11, produced from the combination of the heavy chain CDR3 mutant clone 70-1 with the light chain CDR3. mutant clone 78-34, as having an off rate of 0.0045 ^s-1. This k_{off} rate was a significant improvement compared to the k_{off} rates for either the heavy chain CDR3 mutant clone 70-1 (0.0134 ^s-1), or for the light chain CDR3 mutant clone 78-34 (0.0164 ^s-1) alone. Furthermore, clone 101-11 showed a significant improvement in neutralization assays. Accordingly, clone 101-11 was selected for affinity maturation as described below.

D. Affinity maturation of clone 101-11 consisted of repeat cycles of PCR

mutagenesis of both the heavy and light chain CDR3s of 101-11 using spiked oligonucleotide primers. The clones were selected with decreasing concentrations of biotinylated IL-12 (bio-IL-12). The binding characteristics of the mutated clones was assessed by BIAcore binding analysis and RBA, PHA neutralization assays. The k_{off} rates and/or IC₅₀ values for clones 136-9 through 170-25 are presented in Table 2 (see hinding assay and the PHA blast assay. Clone 103-14 also demonstrated a low k_{off} rate, binding assay and the PHA blast assay. Clone 103-14 also demonstrated a low k_{off} rate, and accordingly was selected for further affinity maturation.

25 E. Generation and Selection of Randomized Libraries of Clone 103-14 Light CDR3 The light chain CDR3 of clone 103-14 (QSYDRGFTGSMV (SEQ ID NO: 590))

was systematically randomized in 3 segments using 3 different libraries as outlined below, where X is encoded by a randomized codon of sequence NMS with N being any nucleotide and S being either deoxycytosine or deoxyguanidine.

L3.2= QSYXXXXXXMV (SEQ ID NO: 592)

L3.3=QSYDRGXXXXXX (SEQ ID NO: 593)

or absence of excess free antigen (p40 and p70). The outputs from selections (clones (H3 and L3.1, L3.2 & L3.3) were constructed and subjected to a large variety of H3) of clone 103-14 was not mutated. Four randomized libraries based on clone 103-14 L3.2, and L3.3) of clone 103-14 was performed. The heavy chain CDR3 (referred to as Randomized mutagenesis of all three light chain CDRs (referred to as L3.1,

73-B1 through 99-G11) were screened primarily by BIAcore, and on occasion with RBA selection conditions that involved using limiting antigen concentration and the presence

Random mutagenesis of the light chain CDR of 103-14 generated clone Y61, and are shown in Table 2 (see Appendix A).

heterodimer. The full length sequences of Y61 heavy chain variable region and light cross-react with Y61 anti-IL-12 antibody to thereby decrease the antibody binding to the not affected by a 50 fold molar excess of free p40, demonstrating that free p40 did not ${\rm IC_{50}}$ value of approximately 130 pM determined by the PHA assay. The ${\rm IC_{50}}$ value was 103-14. Yol was selected for conversion to a whole IgU1. Whole Y61-IgU1 has an which exhibited a significant improvement in IC50 value compared to the parent clone

chain variable region are shown below.

Y61 Heavy Chain Variable Region Peptide Sequence

CDK HI

ОЛОГЛЕЗОССИЛОЬСИЗГИГЗСЧУЗЕЦЕЗ ЗАСМН МЛИОЧЬСКСГЕМЛУ

EIBADC?UKAAVD?AKC BELI?BDN?KNLLYLQMNSLRAEDTAVYYCKT CDK H7 52

HC2HDN MCGCLWALASS (SEG ID NO: 53) срв нз

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Y61 Light Chain Variable Region Peptide Sequence

- 171 -

ÓSAFLÓBBSASCYBCÓKALISC SCCKSNICSNLAK MAGÓFBCLYBKFFIA CDKFI

CDR L2CDR L2

GDK L3 CDK L3 CDK L3

10 CDR residues are assigned according to the Kabat definitions.

EXAMPLE 2: Mutation of Y61 at Hypermutation and Contact Positions

Typically selection of recombinant antibodies with improved affinities can be

contried out using phage display methods. This is accomplished by randomly mutating combinations of CDR residues to generate large libraries containing single-chain antibodies of different sequences. Typically, antibodies with improved affinities are selected based on their ability to reach an equilibrium in an antibody-antigen reaction. However, when Y61 scFV was expressed on phage surface and incubated with IL-12, antibody-antigen equilibrium. The scFV-phage remained bound to IL-12, presumably due to a non-specific interaction, since purified Y61 scFv exhibits normal dissociation due to a non-specific interaction, since purified Y61 scFv exhibits normal dissociation be library generation and selections by mutagenesis of multiple CDR residues) could not be library generation and selections by mutagenesis of multiple CDR positions were mutated utilized, a new strategy involves selection of appropriate CDR positions were mutated.

This strategy involves selection of appropriate CDR positions for mutation and is

based on identification and selection of amino acids that are preferred selective mutagenesis positions, contact positions, and/or hypermutation positions. Contact positions are defined as residues that have a high probability of contact with an antigen when the antigen interacts with the antibody, while hypermutation positions are defined as residues considered to have a high probability for somatic hypermutation during in vivo affinity maturation of the antibody. Preferred selective mutagenesis positions are vivo affinity maturation of the antibody. Preferred selective mutagenesis positions are

was already optimized in the CDR3 regions using the procedure described in Example 1, therefore it was difficult to further improve the area which lies at the center of the antibody binding site using phage-display selection methods. Greater improvements in activity were obtained by mutation of potential contact positions outside the CDR3 regions by either removing a detrimental antigen-antibody contact or, engineering a new

contact.

Amino acids residues of Y61 which were considered contact points with antigen,

and those CDR positions which are sites of somatic hypermutations during in vivo affinity maturation, are shown in Table 3 (see Appendix A). For Y61 affinity maturation, 15 residues outside CDR3, 3 residues within the L3 loop, and 5 residues in

the H3 loop were selected for PCR mutagenesis.

Y61 scFv gene was cloned into the pUC119(Sfi) plasmid vector for mutagenesis.

Oligonucleotides were designed and synthesized with randomized codons to mutate each selected position. Following PCR mutagenesis, a small number of clones (~24) were host cell. The expressed in a host cell, for example, in a bacterial, yeast or mammalian host cell. The expressed antibody was purified and the k_{off} measured using the BIAcore system. Clones with improved off-rates, as compared to Y61, were then tested in neutralization assays. This procedure was repeated for other CDR positions. Individual mutations shown to have improved neutralization activity were combined to generate an antibody with even greater neutralization potency.

The Y61 CDR positions that were mutated in order to improve neutralization potency, and the respective amino-acid substitutions at each position are shown in Figures 2A-2H. Off-rates, as determined by BIAcore analysis, are given. These off rates are also shown in the histograms to the right of each table.

Results of these substitutions at positions H30, H32, H33, H50, H53, H54, H58,

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H95, H97, H101, L50, L92, L93, demonstrated that all amino-acid substitutions examined resulted in antibodies with poorer off-rates than Y61. At positions H52, L32, and L50, only a one amino acid substitution was found to improve the off-rate of Y61, all other changes adversely affected activity. For L50, this single Gly→Tyr change significantly (5-10 times) improved the neutralization potency of Y61. The results demonstrated the importance of these positions to Y61 activity, and suggest that in most cases phage-display was able to select for the optimal residues. However, at positions

- 143 -

H31, H36, L30, and L94, several substitutions were found to improve Y61 off-rate, suggesting that these positions were also important for antigen binding, although the phage display approach did not allow selection of the optimal residues.

Selective mutation of contact and hypermutation positions of Y61 identified amino acid residue L50 in the light chain CDR2, and residue L94 of the light chain CDR3, which improved the neutralization ability of Y61. A combination of these

CDR3, which improved the neutralization ability of Y61. A combination of these mutations produced an additive effect, generating an antibody, 1695, that exhibited a significant increase in neutralization ability. The full length sequence of 1695 heavy and light chain variable region sequences is shown below.

1695 Heavy Chain Variable Region Peptide Sequence

ONOFINESCECONNOLOGISTERSCHASCELLES SYGMH WURQAPGKGLEWVA

EIBADCSUKAAVDSAKC BELISBDNSKNITATÓWNSTBVEDIVAAACKI
CDB H7

TO HC2HDN MCGCLWALASS (SEGIDNO: 31)

CDB H3

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1695 Light Chain Variable Region Peptide Sequence

OSAFLÓBBSASEVBEGKALISC SESKSNICSULAK MAGGFBELVBKFFIA

CDK FJ

KNDÓKЬ ON DE BEROZK SCI SY STYLLOT Ó VEDEYDANC CDK LY

CDK F3 CDK F3

- 14t -

CDR residues are assigned according to the Kabat definitions.

A summary of the heavy and light chain variable region sequence alignments showing the lineage development of clones that were on the path from Joe9 to J695 is shown in Figures 1A-1D. The CDRs and residue numbering are according to Kabat.

EXAMPLE 3: Functional Activity of Anti-hIL-12 Antibodies

To examine the functional activity of the human anti-human IL-12 antibodies of the invention, the antibodies were used in several assays that measure the ability of an antibody to inhibit IL-12 activity.

A. Preparation of Human PHA-activated Lymphoblasts

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Human peripheral blood mononuclear cells (PBMC) were isolated from a leukopac collected from a healthy donor by Ficoll-Hypaque gradient centrifugation for 45 minutes at 1500 rpm as described in Current Protocols in Immunology, Unit 7.1. PBMC at the interface of the aqueous blood solution and the lymphocyte separation medium were collected and washed three times with phosphate-buffered saline (PBS) by centrifugation for 15 minutes at 1500 rpm to remove Ficoll-Paque particles.

The PBMC were then activated to form lymphoblasts as described in Current

Protocols in Immunology, Unit 6.16. The washed PBMC were resuspended at 0.5-1x106 cells/ml in RPMI complete medium (RPMI 1640 medium, 10% fetal bovine serum (FBS), 100 U/ml penicillin, 100 µg/ml streptomycin), supplemented with 0.2% (v/v) PHA-P (Difco, Detroit, MI) and cultured for four days at 37°C in a 5% CO₂ atmosphere.

After four days, cell cultures were split 1:1 by volume in RPMI complete medium, plus 0.2% (v/v) PHA-P and 50 U/ml recombinant human IL-2. Recombinant human IL-2 was produced by transfection of an expression vector carrying the human IL-2 cDNA into COS cells (see Kaufman et al., (1991) Nucleic Acids Res. 19, 4484-4490), and purified as described in PCT/US96/01382. Cell cultures were then incubated for an additional one to three days. PHA blast cells were harvested, washed twice with RPMI additional one to three days. PHA blast cells were harvested, washed twice with RPMI complete medium and frozen in 95% FBS, 5% DMSO at 10x10⁶ cells/ml.

PHA blast cells to be used for the IL-12 receptor binding assay (see section B) were collected after one day culture in the presence of IL-2, whereas PHA blast cells to be used for the PHA blast proliferation assay (see section C) and the interferon-gamma induction assay (see section D) were collected after three day culture in the presence of IL-2.

B. IL-12 Receptor Binding Assay

Incubations were carried out in duplicate.

The ability of anti-IL-12 antibodies to inhibit binding of radiolabelled IL-12 to antibodies to inhibit binding of radiolabelled IL-12 to antibody were analyzed as follows. Various concentrations of anti-IL-12 antibody were preincubated for 1 hour at 37°C with 50-100 pM ¹²⁵I-hIL-12 (iodinated hIL-12 was prepared using the Bolton-Hunter labeling method to a specific activity of 20-40mCi/mg from MEN-Dupont) in binding buffer (RPMI 1640, 5% FBS, 25 mM Hepes pH 7.4). PHA blast cells isolated as described above, were washed once and resuspended in binding buffer to a cell density of 2x10⁷ cells/ml. PHA blasts (1x10⁶ cells) were added to the antibody ¹²⁵I-hIL-12 mixture and incubated for two hours at room temperature. Cell bound radioactivity was separated from free ¹²⁵I-hIL-12 by centrifugation of the assay mixture for 30 seconds at room temperature, aspiration of the liquid and a wash with 0.1ml binding buffer, followed by centrifugation at 4°C for 4 min at 10,000 x g. The cell pellet was examined for cell bound radioactivity using a gamma counter. Total binding was determined in the absence of antibody and non-specific binding was determined by inclusion of 25 nM unlabeled IL-12 in the assay.

In the IL-12 receptor binding assay using the Y61 and J695 human anti-IL-12 antibodies, both antibodies demonstrated a comparable inhibition of IL-12 receptor binding with an IC₅₀ value of approximately 1.6 x 10⁻¹¹M, while J695 had an IC₅₀ value of approximately 1.1 x 10⁻¹¹M.

Anti-IL-12 antibodies were evaluated for their ability to inhibit PHA blast

C. Human PHA Blast Proliferation Assay

proliferation (which proliferation is stimulated by IL-12). Setial dilutions of anti-IL-12 antibody were preincubated for 1 hour at 37°C, 5% CO₂ with 230 pg/ml hIL-12 in 100 ml RPMI complete medium in a microtiter plate (U-bottom, 96-well, Costar, resuspended in RPMI complete medium to a cell density of 3x10⁵ cells/ml. PHA blasts (100 ml, 3x10⁴ cells) were added to the antibody/hIL-12 mixture, incubated for 3 days at 37°C, 5% CO₂ and labeled for 4-6 hours with 0.5 mCi/well (3H)-Thymidine (Amersham, Arlington Heights, IL). The culture contents were harvested onto glass fiber filters by means of a cell harvester (Tomtec, Orange, CT) and (³H)-Thymidine

samples were assayed in duplicate. The results of neutralization in the presence of varying concentrations of p70:p40

incorporation into cellular DNA was measured by liquid scintillation counting. All

5 (i.e. the ratio of IL-12 heterodimer to free p40 subunit) is shown in Table 4 (see

Appendix A).

Analysis of the Y61 human anti-IL-12 antibody in the PHA blast proliferation assay demonstrated that the antibody inhibited PHA blast proliferation with an IC₅₀

value of approximately 1.8 x 10⁻¹⁰ M in the presence of IL-12 p70 alone, without any

excess p40 (p70:p40 ratio of 1:0). In the presence of a 50-fold excess of free p40 (p70:p40 at a ratio of 1:50), the Y61 antibody inhibited PHA blast proliferation with an IC₅₀ value of approximately 1.8 x 10⁻¹⁰M. This result demonstrates that the ability of Y61 to inhibit blast proliferation is not compromised by the presence of excess p40. The human anti-IL-12 antibody, J695 inhibited PHA blast proliferation with an

IC₅₀ value of approximately 1.0 x 10⁻¹¹M in the presence of p70:p40 at a ratio of 1:0. In the presence of a p70:p40 ratio of 1:50, this antibody inhibited PHA blast proliferation with an IC₅₀ value of approximately 5.8 ± 2.8 x 10⁻¹² M (n=2), demonstrating that the excess p40 had only a slight inhibitory effect on the antibody. Overall results demonstrate the improved neutralization activity of J695 in comparison with Y61 due to the mutations at L50 and L94.

D. Interferon-gamma Induction Assay

The ability of anti-IL-12 antibodies to inhibit the production of IFNy by PHA blasts (which production is stimulated by IL-12) was analyzed as follows. Various concentrations of anti-IL-12 antibody were preincubated for 1 hour at 37°C, 5% CO₂ with 200-400 pg/ml hIL-12 in 100 ml RPMI complete medium in a microtiter plate once and resuspended in RPMI complete medium to a cell density of 1x10⁷ cells/ml. PHA blasts (100 µl of 1x10⁶ cells) were added to the antibody/hIL-12 mixture and incubated for 18 hours at 37°C and 5% CO₂. After incubation, 150 µl of cell free incubated for 18 hours at 37°C and 5% CO₂. After incubation, 150 µl of cell free incubated for 18 hours at 37°C and 5% CO₂. After incubation, 150 µl of cell free incubated for 18 hours at 31°C and 5% CO₂. After incubation, 150 µl of cell free incubated for 18 hours at 31°C and 5% CO₂. After incubation, 150 µl of cell free incubated for 18 hours at 31°C and 5% CO₂. After incubation, 150 µl of cell free incubated for 18 hours at 31°C and 5% CO₂. After incubation, 150 µl of cell free incubated for 18 hours at 31°C and 5% CO₂. After incubation, 150 µl of cell free incubated for 18 hours at 31°C and 5% CO₂. After incubation, 150 µl of cell free incubated for 18 hours at 31°C and 5% CO₂. After incubation, 150 µl of cell free incubated for 18 hours at 31°C and 5% CO₂. After incubation, 150 µl of cell free incubated for 18 hours at 31°C and 5% CO₂. After incubation with an IC₅₀ value of approximated that

while the human anti-IL-12 antibody, 1695, inhibited human IFNy production with an

IC₅₀ value of approximately $5.0 \pm 2.3 \times 10^{-12} \,\mathrm{M}$ (n=3). The result demonstrates the substantial improvement in the affinity of J695 as a result of the modifications at L50 and L94.

20 E. Induction of Non-human IL-12 from Isolated PBMC

To examine the cross-reactivity of the human anti-hIL-12 antibodies with IL-12 from other species, non-human IL-12 was produced as follows. PBMC were separated from fresh heparinized blood by density gradient centrifugation as described above using lymphoprep (Mycomed, Oslo, Norway) for cynomolgus monkey, baboon, and dog, PBMC, Accu-paque (Accurate Chemical & Sci. Corp., Westbury, NY) for dog PBMC or Lympholyte-rat (Accurate Chemical & Sci. Corp., Westbury, NY) for rat PBMC.

The PBMC were then induced to produce IL-12 as described (D'Andrea et al., (1992) J.Exp. Med 176, 1387-1398, Villinger et al., (1995) J. Immunol. 155, 3946-

3954, Buettner et al., (1998) Cytokine 10, 241-248). The washed PBMC were resuspended at 1x106 cells/ml in RPMI complete medium, supplemented with 0.0075% (wt/vol) of SAC (Pansorbin; Calbiochem-Behring Co., La Jolla, CA) or 1-5 mg/ml

ConA (Sigma Chemical Co., St. Louis, MO) plus 0.0075% SAC and incubated for 18 hours at 37°C in a 5% CO₂ atmosphere. Cell-free and SAC-free medium was collected by centrifugation and filtering through 0.2 mm filters.

IL-12 from the rhesus monkey was obtained as recombinant thesus IL-12 from

5 Emory University School of Medicine, Atlanta, GA.

F. Murine 2D6 Cell Proliferation Assay

The murine T cell clone 2D6 proliferates in response to murine IL-2, IL-4, IL-7

and IL-12 (Maruo et al., (1997) J. Leukocyte Biol. 61, 346-352). A significant proliferation was also detected in response to rat PBMC supernatants containing rat IL-12. The cells do not respond to dog, cynomolgus, baboon or human IL-12. Murine 2D6 cells were propagated in RPMI complete medium supplemented with 50 mM betamercaptoethanol (BME) and 30 ng/ml murine IL-12. One day prior to the assay, the murine IL-12 was washed out and the cells were incubated overnight in RPMI complete medium plus BME.

Serial dilutions of anti-IL-12 antibody were preincubated for 1 hour at 37°C, 5% CO₂ with 40 pg/ml murine IL-12 in 100 ml RPMI complete medium plus βME in a microtiter plate (U-bottom, 96-well, Costar). 2D6 cells were washed once and resuspended in RPMI complete medium containiny βME to a cell density of 1x10⁵ cells/ml. 2D6 cells (100 µl, 1x10⁴ cells) were added to the antibody/hIL-12 mixture, incubated for 3 days at 37°C, 5% CO₂ and labeled for 4-6 hours with 0.5 mCi/well (3H)-recubated for 3 days at 37°C, 5% CO₂ and labeled and counted by liquid scintillation counting. All samples were assayed in duplicate.

25 G. Species Cross-reactivity of 1695 with Non-Human IL-12 was analysed

Species cross-reactivity of J695 with non-human IL-12 was analyzed using PBMC's isolated from several non-human species. The presence of non-human IL-12 activity in the rat, dog, cynomolgus and baboon PBMC supernatants was confirmed using several bioassays described above, such as the murine 2D6 cell proliferation assay, the human PHA blast proliferation assay and the interferon-gamma induction assay by blocking the non-human PBMC induced responses with rabbit and/or sheep polyclonal

antibodies Y61 and J695 with non-human IL-12 in PBMC supernatants or purified murine and rhesus IL-12 was then assessed in the same bioassay(s) by determining the J695 antibody concentration at which 50% inhibition of the response was observed. The species cross-reactivity results are summarized in Table 5. The results demonstrate that Y61 and J695 are each able to recognize IL-12 from monkeys (e.g., cynomolgus and rhesus IL-12 for Y61, and cynomolgus, rhesus and baboon for J695) and that J695 is approximately 35 fold less active on dog IL-12; neither Y61 nor J695 cross reacts with mouse or rat IL-12.

H. Human cytokine specificity of 1695

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The specificity of J695 was tested in a competition ELISA in which a panel of human cytokines was tested for their ability to interfere with the binding of soluble J695 to immobilized human IL-12. The panel of human cytokines included IL-1α and IL-1β (Genzyme, Boston, MA), IL-2 (Endogen), IL-4, IL-10, IL-17, IFN-gamma, and TGF-β1 (R&D, Minneapolis, MA) IL-8 (Calbiochem), PDGF, IGF-I, and IGF-II (Boehringer Mannheim Corp., Indianapolis, IN), TMFα and lymphotoxin, IL-6, soluble IL-6 receptor, IL-11, IL-12 p70, IL-12 p40, M-CSF, and LIF. EBI-3, an IL-12 p40 related protein that is induced by Epstein-Barr virus infection in B lymphocytes (Devergne et protein that is induced by Epstein-Barr virus infection in B lymphocytes (Devergne et

Antibody human-αhull12 mouse-ahull 12 rabbit-amuIL12 rat-omult 12 Specificity Non-neutralizing Mouse IL-12 3.0 x 10⁻¹¹ 1.5 x 10⁻¹⁰ Purified PBMC sup 6.0 x 10⁻¹⁰ Rat IL-12 Dog IL-12 IC₅₀ (M) PBMC sup 2.2 x 10⁻¹⁰ Cyno IL-12 1.2 x 10⁻¹⁰ PBMC sup Rhesus IL-12 1.0 x 10-10 1.0×10^{-10} Purified Baboon IL-12 PBMC sup 2.0 x 10⁻¹⁰

C17.15

761 1695

human-ahull 12

Non-neutralizing

Non-neutralizing

3.5 x 10⁻¹⁰

1.0 x 10-11

1.0 x 10⁻¹¹

1.5 x 10⁻¹¹

1.7 x 10⁻¹⁰

5.0 x 10⁻¹¹

Name

Table 5 Species Cross Reactivity Data

Human IL-1

Purified

al., (1996) J. Virol. 70, 1143-1153) was expressed as a human lgG-Fc chimera (EBI-3/Fc) Single-stranded salmon sperm DNA (Sigma) was also tested.

Flat-bottom ELISA immunoassay microtiter plates (96 well, high binding,

Costat) were coated overnight at 4°C with 0.1 ml human IL-12 (2 µg/ml in 0.1 M NaHCO₃)).

5 carbonate coating buffer (4 volumes 0.1 M NaHCO₃ plus 8.5 volumes 0.1 M NaHCO₃)).

The plates were washed twice with PBS containing 0.05 % Tween 20 (PBS-T), blocked with 200µl of 1 mg/ml bovine serum albumin (BSA, Sigma) in PBS-T for 1 hour at 1L-12 antibody 1695 (100 ng/ml) and each cytokine (2nM) in PBS-T containing 50 plates were washed 4 times and incubated for 2 h at room temperature. The plates were washed 4 times and incubated for 1h at room temperature with 100 µl mouse anti-human lambda-HRP (1:500 in PBS-T/BSA, Southern Biotech. Ass. Inc., Birmingham, AL). The plates were washed 4 times and developed with ABTS

(Kirkegaard & Perry Lab., Gaithersburg, MD) for 20-30 minutes in the dark. The OD₄₅₀nm was read using a microplate reader (Molecular Devices, Menlo Park, CA).

Percent binding was determined relative to 1695 binding to the IL-12 coated plate in the absence of any soluble cytokine.

blocked only by human IL-12 p70 and to a lesser extent, by human IL-12 p40 and not by

The results demonstrated that 1695 binding to immobilized human IL-12 was

I. Binding to a Novel IL-12 Molecule

any of the other cytokines tested.

An alternative IL-12 heterodimer has been described, in which the p35 subunit is replaced by a novel p19 molecule. P19 was identified using 3D homology searching for IL-6/IL-12 family members, and is synthesized by activated dendritic cells. P19 binds to p40 to form a p19/p40 dimer, which has IL-12 –like activity, but is not as potent as the p35/p40 heterodimer in IFNy induction. Antibodies which recognize p40 alone, but preferably in the context of a p70 molecule (e.g., 1695 and Y61, see Example 3H) are expected to also neutralize both the p35/p40 molecules and the p19/p40 molecules.

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In vivo Activity of Anti-hIL-12 Antibodies EXYMbre 4:

decrease in the platelet count was observed at 72 hours. Levels of plasma neopterin, a cell count (WBC), especially in the lymphocyte and monocyte subsets after 24 hours. A human IL-12 at 1 µg/kg/day for a period of 5 days resulted in a decrease in white blood Biophys Res. Comm. 204: 1150-1157. In those previous studies, administration of IL-12 on peripheral hematology in cynomolgus monkey Bree et al., (1994) Biochem examined in a model modified from one used by Bree et al. to study the effect of human The in vivo effects of IL-12 antibodies on IL-12 induced responses were

In the first study with human anti-hIL-12 antibodies, fifteen healthy cynomolgus were the highest at 72 hours. marker of monocyte activation in response to IFN-y, began to elevate at 24 hours and

10 mg/kg C8.6.2. Group 4 received an intravenous administration of 1 mg/kg Y61 human IL-12 monoclonal antibody). Group 3 received an intravenous administration of 2 received an intravenous administration of 1 mg/kg C8.6.2 (neutralizing mouse antiimmunoglobulin (IVIG, Miles, Eckhart, IN, purified using protein A Sepharose). Group Group 1 received an intravenous (IV) administration of 10 mg/kg human intravenous monkeys with an average weight of 5kg, were sedated and divided into 5 groups (n=3).

Group 5 received an intravenous administration of 10 mg/kg Y61. (human anti-human IL-12 antibody, purified from CHO cell conditioned medium).

complete blood cell counts with differentials and serum chemistry. Serum human IL-12, the following time points: baseline, 8, 24, 48, 96 and 216 hours, and analyzed for subcutaneous (SC) injection of human IL-12 (1 µg/kg). Blood samples were taken at One hour after the antibody administration all animals received a single

Animals treated with IL-12 plus IVIG control antibody (Group 1) showed many plasma neopterin levels were also measured. C8.6.2 antibody, Y61 antibody, monkey IFN-gamma, monkey IL-10, monkey IL-6 and

pronounced in the animals treated with either the C8.6.2 or Y61 antibody at 1 or 10 lymphocyte count and monocyte count. These decreases were not seen or were less of the expected hematological changes, including decreases in WBC, platelets,

mg/kg (Groups 2-5).

Serum or plasma samples were analyzed by ELISA specific for monkey IFN-gamma and monkey IL-10 (Biosource International, Camarillo, CA), monkey IL-6 (Endogen) and plasma neopterin (ICM Pharmaceuticals. Orangeburg, NY). IFM-gamma, IL-10 or IL-6 were not detected in any of the IL-12 treated animals including the control animals treated with IL-12 plus IVIG. This was probably due to the low level exposure to IL-12 (only 1 dose of 1 µg/kg). Nevertheless, plasma neopterin levels increased

to IL-12 (only 1 dose of 1 µg/kg). Nevertheless, plasma neopterin levels increased about three fold in the IL-12 plus IVIG treated animals but did not change in all C8.6.2 or Y61 treated animals, including the lower dose (1 mg/kg) Y61 treated animals, indicating that Y61 was effective in vivo in blocking this sensitive response to IL-12. In a second study, in vivo activity and pharmacodynamics (PD) of J695 in

cynomolgous monkeys were studied by administering exogenous rhlL-12 and determining if 1695 could block or reduce the responses normally associated with rhlL-12 and 12 administration. Male cynomolgus monkeys (n=3 per group) were administered a single dose of 0.05, 0.2, or 1.0 mg/kg 1695 or 1 mg/kg intravenous immunoglobulin (SC) in the dorsal skin. One hour following the administration of 1695 or IVIG, all animals received a single SC dose of 1 µg/kg rhlL-12 in the dorsal skin. Blood samples animals received a single SC dose of 1 µg/kg rhlL-12 in the dorsal skin. Blood samples are collected via the femoral vein up to 28 days after 1695 administration. Serum was acquired from each blood sample and assayed for IL-12, 1695, IFN-γ, and anti-1695 acquired from each blood sample and assayed by reverse-phase high performance liquid antibodies by ELISA. Neopterin was assayed by reverse-phase high performance liquid

chromatography. The levels of neopterin, normalized with respect to the levels of neopterin that

were measured before administration of J695 or rhIL-12, are shown in Figure 3. To

normalized for neopterin levels was calculated for each animal (Table 6). Neopterin exposure (AUC) was suppressed in a dose-dependent manner between approximately 71 and 93% in the IV groups and between 71 and 100% in SC groups, relative to the IVIG control groups. These results suggest that the dose of 1695 necessary for 50% inhibition of the neopterin response (ED₅₀) was less than 0.05 mg/kg when administered by either

compare the suppression of neopterin between groups, the area under the curve (AUC)

the IV or SC route.

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Table 6: Dose-Dependent Suppression of IL-12 Induced Neopierin by 1695 in Cynomolgus Monkeys

100	601 ∓ 0	_	0.1	DS nəvig 21-11 namud
£.£7	6'St ∓ S6E	-	2.0	py a dose of 1 µg/kg
7.17	450 ∓ 108	-	č0.0	followed I hour later
0	t09 = 08t!	0.1	-	noitosini DS əlgnis
L.26	158 ∓ 565	-	0.1	DS nevig 21-JI
9.88	91€ ∓ 661	<u>-</u>	2.0	qoze ot Ihg/kg human
£.17	205 ∓ 135	-	80.0	followed I hr later by a
00	St8 ∓ St/1	0.1	-	noisoəjni VI əlgni2
Control				
Compared with	Neopterin Levels			
Neopterin AUC	Normalized	(ឱង/ឱពា)	(ឃឥ/ុ៩ឥ)	or J695 and rhlL-12
% Reduction of	NO OUA	IAIC Dose	Jegs Dose	Noute of dosing IVIG

Treatment with 1695 also prevented or reduced the changes in hematology normally associated with rhIL-12 administration (leukopenia and thrombocytopenia). At 24 hours after rhIL-12 administration lymphocyte counts were reduced by approximately 50% when compared to baseline values in the control IV and SC IVIG treated groups. Administration of 1695 either SC or IV at all three dose levels prevented this reduction, resulting in lymphocyte counts at 24 hours approximately the same as baseline values. At 48 hours after IL-12 administration, platelet counts in the groups treated with IV and SC IVIG were reduced by approximately 25% when compared to baseline values.

An example dose schedule targeted to maintain serum levels above the 90% effect level would be 1 mg/kg IV and SC given approximately every other week, or 0.3 mg/kg given approximately every week, assuming slight accumulation during repeated dosing. This study demonstrates that antibody can be given safely to monkeys at such dosages. In independent toxicity studies, it was further found that up to 100 mg/kg of the antibody can be given safely to monkeys.

chimeric IL-12, a molecule which combines the murine p35 subunit with the human IL-12 p40 subunit. In contrast to human IL-12 which is biologically inactive in mice, this

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J695 was also effective in preventing IFN-y production in mice treated with a

chimeric IL-12 retains biological function in mice, including induction of IFN-γ. In addition, the human p40 subunit allows the molecule to be bound and neutralized by 1695. Chimeric IL-12 at a dose of 0.05 mg/kg i.p. was administered to female C3H/He1 mice (10/experimental group) in five daily doses on days 0, 1, 2, 3, and 4. 1695 was given on days 0, 2 and 4 at doses of 0.05, 0.01, 0.002, 0.0004, 0.00008, and 0.000016 mg/kg i.p., 30' prior to the IL-12 injections. The control hulgG1γ was given IP, at a dose of 0.05 mg/kg on days 0, 2, and 4. The mice were bled on day 5, and serum IFN-γ levels were determined by ELISA. The results demonstrated that 1695 caused dose-levels were determined by ELISA. The results demonstrated that 1695 caused dose-levels were determined by ELISA. The results demonstrated that 1695 caused dose-levels were determined by ELISA. The results demonstrated that 1695 caused dose-levels were determined by ELISA. The results demonstrated that 1695 caused dose-levels were determined by ELISA is a potent inhibitor of II-12 activity in vivo.

EXAMPLE 5: Kinetic Analysis of Binding of Human Antibodies to Recombinant human IL-12 (rhIL-12)

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antibody 1695, captured on a biosensor matrix) and analyte (rhIL12 in solution) were measured by surface plasmon resonance (SPR) using the BIAcore system (Biacore AB, Uppsala, Sweden). The system utilizes the optical properties of SPR to detect covalently bound to the dextran matrix at known concentrations. Antibodies are injected immobilized ligand results in an increased matrix protein concentration and resultant change in the SPR signal. These changes in SPR signal are recorded as resonance units change in the SPR signal. These changes in SPR signal are recorded as resonance units change in the SPR signal. These changes in SPR signal are recorded as resonance units change in the SPR signal. These changes in SPR signal are recorded as resonance units change in the SPR signal are recorded as resonance units change in the SPR signal are recorded as resonance units change in the SPR signal are recorded as resonance units change in the SPR signal are recorded as resonance units change in the SPR signal are recorded as resonance units change in the SPR signal are recorded as resonance units change in the SPR signal are recorded as resonance units change in the SPR signal are recorded as resonance units change in the SPR signal are recorded as resonance units change in the SPR signal are recorded as resonance units change in the SPR signal are recorded as resonance units change in the SPR signal are resonance units are specific plants.

Real-time binding interactions between captured ligand (human anti-rhIL-12

Associates, Cat. No. 2040-01, Birmingham, AL) on the biosensor matrix, goat anti-human IgG is covalently linked via free amine groups to the dextran matrix by first activating carboxyl groups on the matrix with 100 mM N-hydroxysuccinimide (NHS) and 400 mM N-Ethyl-N'-(3-dimethylaminopropyl)-carbodiimide hydrochloride (EDC). Next, goat anti-human IgG is injected across the activated matrix. Thirty-five microliters of goat anti-human IgG (25 µg/ml), diluted in sodium acetate, pH 4.5, is

injected across the activated biosensor and free amines on the protein are bound directly to the activated carboxyl groups. Unreacted matrix EDC-esters are deactivated by an injection of 1 M ethanolamine. Standard amine coupling kits were commercially available (Biacore AB, Cat. No. BR-1000-50, Uppsala, Sweden).

1695 was diluted in HBS running buffer (Biacore AB, Cat. No. BR-1001-88,

Representative results of CHO derived J695 binding to rhIL-12 as compared to BIAcore kinetic evaluation software (version 2.1) was used. sample. To determine the dissociation constant (off-rate), association constant (on-rate), Biosensor matrices were regenerated using 100 mM HCI before injection of the next of cytokine injection was taken to represent the binding value of the particular sample. through each flow cell. The net difference in baseline signal and signal after completion Prior to injection of thIL-12, and immediately afterwards, HBS buffer alone flowed concentrations of rhlL-12 employed were 10, 20, 25, 40, 50, 80, 100, 150 and 200 nM. injected through the immobilized protein matrices at a flow rate of 5μ /min. The was measured. Cytokines were diluted in HBS running buffer and 50 µl aliquots were (approximately 1200 RU's). Direct rhlL12 specific antibody binding to soluble rhlL12 completion of 1695 injection was taken to represent the amount of 1gG1 J695 bound between the baseline and the point corresponding to approximately 30 seconds after afterward, HBS buffer alone flowed through each flow cell. The net difference in signal matrix at a flow rate of 5 µl/min. Before injection of the protein and immediately were injected through the goat anti-human IgO polyclonal antibody coupled dextran binding assay was conducted as follows. Aliquots of 1695 (25 µg/ml; 25 µl aliquots) the capacity of rhlL12-specific antibodies to bind immobilized goat anti-human lgG, a Uppsala, Sweden) to be captured on the matrix via goat anti-human IgG. To determine

Table 7: Binding of CHO or COS derived J695 to thIL-12.

the COS derived J695, are shown in Table 7.

£4.1	1490	766	100	СНО
24.1	1272	1033	051	СНО
84.1	191	1112	200	СНО
Thili2/AB	Ab, bound, RU's	rhlL12 bound, RU's	Mn.,SLIIh	Source

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00.1	LLEI	779	01	COS
07.1	1394	£8 <i>L</i>	50	cos
	601	833	52_	COS
1.34	1431	b 68	07	cos
7£.1	1457	726	0\$	cos
74.1	1486	586	08	SOO
1.44	1254	1024	100	cos
94.1	9851	108⊄	0\$1	cos
67.1	0691	1115	500	cos
HIL12/AB	s'U's, bnuod, dA	rhlL12 bound, RU's	Mn.121.11h	Source
86.0	1251	<i>L</i> 79	01	СНО
07.1	1382	£ <i>LL</i>	50	СНО
1.25	1398	818	52	СНО
55.1	1413	LL8	. 07	СНО
95.1	1434	615	05	СНО
1.40	LSti	\$\$6	08	СНО

Molecular kinetic interactions between captured J695 and soluble rhlL-12 were quantitatively analyzed using BIAcore technology. Several independent experiments were performed and the results were analyzed by the available BIAcore mathematical analysis software to derive kinetic rate constants, as shown in Table 8.

Table 8: Apparent kinetic rate and affinity constants of 1695 for rhIL-12.

9.74E-11	7.61E-05	3.40E+05	cos	5691
1.34E-10	4'72E-05	3'25E+02	СНО	\$691
.gvA	.gvA	.gvA		
Kq (M)'	Off-rate (s-1),	On-rate (M-1s-1),	Source	Antibody

10

There was a small difference between the calculated apparent constant (Kd) for the interaction between CHO derived 1695 (Kd = $1.34^{-10}M^{-1}$) and COS derived 1695 (Kd = $9.74 \times 10^{-11}M^{-1}$) antibodies. The apparent dissociation constant (Kd) between 1695 and rhIL12 was estimated from the observed rate constants by the formula: Kd = off-tate\) on-rate.

To determine the apparent association and dissociation rate constant for the interaction between J695 and rhIL-12, several binding reactions were performed using a fixed amount of J695 (2 µg/ml) and varying concentrations of rhIL-12. Real-time binding interaction sensorgrams between captured J695 and soluble rhIL12 showed that both forms of antibody were very similar for both the association and dissociation phase. To further evaluate the capacity of captured IgG1 J695 mAb to bind soluble

recombinant cytokine, a direct BIAcore method was used. In this method, goat antihuman IgG (25 µg/ml) coupled carboxymethyl dextran sensor surface was coated with IgG1 1695 (2µg/ml) and recombinant cytokine was then added. When soluble rhIL12 was injected across a biosensor surface captured with CHO or COS derived IgG1 1695, the amount of signal increased as the concentration of cytokine in the solution increased. No binding was observed with rmIL12 (R&D Systems, Cat. No. 419-ML, Minneapolis, Mo binding was observed with rmIL12 (R&D Systems, Cat. No. 419-ML, Minneapolis, on the IL12 any concentration tested up to 1000 nM. These results support the conclusion that IgG1 1695 antibodies recognize a distinct determinant on rhIL-12.

Table 9 shows the results of an experiment using BIAcore to demonstrate human tested with the conclusion that IgG1 1695 antibodies recognize a distinct determinant on rhIL-12.

IgG1 J695 mAb binding to only soluble thIL12 and none of the other recombinant cytokines.

Table 9: Epitope mapping of 1695 using BIAcore technology.

Negative	Negative	rec. murine IL12
Poiiisof	Positive	rec. human IL12
		Soluble analyte
Captured ligand CHO 1695	Captured ligand COS 1695	

EXAMPLE 6: Further Studies of J695 Affinity for IL-12

Molecular kinetic interactions between 1695 antibody and human IL-12 were kinetic rate constants were derived.

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BIAcore technology was used to measure the binding of soluble rhlL-12 to solid

 K_d value for the interaction (see Table 10). The results indicated that J695 has a high dissociation (k_a) and association (k_a) rate constants were derived and used to calculate a are averages for the three experiments. From these measurements, the apparent 1695 and IL-12. Three independent experiments were performed, and the values shown order association kinetics, as well as a simple one-to-one molecular interaction between association rate constants were calculated, assuming zero-order dissociation and first concentrations to 1695 was measured as a function of time. Apparent dissociation and Varying concentrations of rhlL-12 were applied, and the binding of IL-12 at different biosensor chips, then a fixed amount of J695 was injected and captured on the surface. phase captured 1695. A goat anti-human IgG antibody was immobilized on the

affinity for rhlL-12.

Table 10: Kinetic Parameters for the Interaction Between J695 and Human IL-12

(Jm\gn 41) M 11-01 x 47.6	K ^q
$^{1-2}$ ¹ - 1 - $^{$	Κ ^υ
1 ·s 8 ·01 × 04·0 ± 17.8	Κ ^q
-Value	Kinetic Parameter

Monoclonal Antibody to Murine Interleukin-12 EXVMbre 1: Characteristics and Neutralization Activity of C17.15, a Rat

IL-12 binding to cell surface receptors, was assessed, as were the kinetics of the C17.15neutralize murine IL-12 activity in a PHA blast proliferation assay, and to block murine monoclonal antibody with murine IL-12, was examined. The ability of C17.15 to approaches in human disease, the interaction of CI7.15, a rat anti-murine IL-12 and autoimmunity using monoclonal antibodies specific for murine IL-12 to similar To assess the relevance of IL-12 treatment studies in mouse models of inflammation 50

murine IL-12 binding interaction.

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In a human PHA blast proliferation assay (See Example 3), serial dilutions of C17.15 or rat 1gG2a (a control antibody) were preincubated with 230 pg/mL murine 1L-12 for 1 hr at 37 °C. PHA-stimulated blast cells were added to the antibody-IL-12 mixtures and incubated for 3 days at 37 °C. The cells were subsequently labeled for 6 h incorporation was measured. Background non-specific proliferation was measured in the absence of added murine IL-12. All samples were assayed in duplicate. The IC₅₀ (M) of C17.15 for recombinant murine IL-12 in this assay was found to be 1.4 x 10⁻¹¹, as compared to the IC₅₀ value of 5.8x10⁻¹² observed for J695 for recombinant human IL-12 under the same conditions (see Table 11).

Table 11: Comparison of the properties of anti-human IL-12 monoclonal antibody J695 and the rat anti-mouse IL-12 monoclonal antibody C17.15

PHA blast Assay	Receptor Binding Assay	Biomolecular Interaction Assay			Epitope	ybodiinA
IC ₅₀ (M)	IC ⁵⁰ (M)	K ^q (W)	k _d , off-rate (s ⁻¹)	k _a , on-rate (M ⁻¹ s ⁻¹)		
₇₁ .0[x 8.¢	11-01 x 1.1	11-01 × 47.6	c-01 × 17.8	3.81 × 10 ⁵	0⊅q uH	\$69r
TF-01 x 4.1	⁰¹⁻ 01 x č.1	4.80 × 10 ⁻¹⁰	+-01 x 48.1	² 01 x 08.£	04q uM	\$1.715

The ability of C17.15 to inhibit the binding of murine 1L-12 to cellular receptors was also measured. Serial dilutions of C17.15 were pre-incubated for 1 hr at 37 °C with 100 pM [¹²⁵I]-murine 1L-12 in binding buffer. The 2D6 cells (2x10⁶) were added to the antibody/[¹²⁵I]-murine 1L-12 mixture and incubated for 2 hours at room temperature.

Cell-bound radioactivity was separated from free [¹²⁵I]-1L-12, and the remaining cell-bound radioactivity was determined. Total binding of the labeled murine 1L-12 to receptors on 2D6 cells was determined in the absence of antibody, and non-specific binding was determined by the inclusion of 25 nM unlabelled murine 1L-12 in the assay.

Specific binding was calculated as the total binding minus the non-specific binding.

Incubations were carried out in duplicate. The results showed that C17.15 has an IC₅₀ and the incubations of binding of murine IL-12 to cellular receptors.

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The affinity of C17.15 for recombinant murine IL-12 was assessed by biomolecular interaction analysis. A goat anti-rat IgG antibody was immobilized on the biosensor capture of C17.15 on the surface of the chip. Varying concentrations of recombinant immobilized C17.15 on the surface of the chip. Varying concentrations of recombinant association rate constants were calculated, assuming a zero order dissociation and first order association kinetics as well as a simple one to one molecular interaction between order association kinetics as well as a simple one to one molecular interaction between dissociation (k_a, off-rate) and association (k_a, on-rate) rate constants were calculated as a simple one to one molecular interaction between dissociation (k_a, off-rate) and association (k_a, on-rate) rate constants were calculated.

These results were used to calculate a K_a value for the interaction. An on-rate of 3.8 x of 3 M⁻¹s⁻¹, an off-rate of 1.84 x 10⁻⁴ s⁻¹, and a K_a of 4.8 x 10⁻¹⁰ was observed for the recombinant murine IL-12-C17.15 interaction.

to cell surface receptors, as well as the kinetics of binding of C17.15 to murine IL-12 correlate with similar measurements for the 1695-rhIL-12 interaction. This indicates that the modes of action of the rat anti-mouse IL-12 antibody C17.15 and anti-human IL-12 antibody 1695 are nearly identical based upon on-rate, off-rate, K_d, IC₅₀, and the PHA blast assay. Therefore, C17.15 was used as a homologous antibody to 1695 in murine models of inflammation and autoimmune disease to study the effects of IL-12 blockade on the initiation or progression of disease in these model animals (see Example blockade on the initiation or progression of disease in these model animals (see Example blockade on the initiation or progression of disease in these model animals (see Example blockade on the initiation or progression of disease in these model animals (see Example blockade on the initiation or progression of disease in these model animals (see Example blockade on the initiation or progression of disease in these model animals (see Example blockade on the initiation or progression of disease in these models of initiation or progression of disease in these models of initiation or progression of disease in these models are initiation or progression of disease in the land of the land of the land of the land of the land of the land of the land of the land of the land of the land of the land of the land of land o

EXAMPLE 8: Treatment of Autoimmune or Inflammation-Based Diseases in Mice by a-Murine IL-12 Antibody Administration

A. Suppression of Collagen-Induced Arthrits in Mice by the α-II-12 antibody CI7.15
 A correlation between IL-12 levels and rheumatoid arthritis (RA) has been demonstrated. For example, elevated levels of IL-12 p70 have been detected in the synovia of RA patients compared with healthy controls (Morita et al (1998) Arthritis and Rheumatism. 41: 306-314). Therefore, the ability of CI7.15, a rat anti-mouse IL-12 and theumatism of Rathritis in mice was assessed.

Male DBA/1 mice (10/group) were immunized with type II collagen on Day 0 and treated with C17.15, or control rat IgG. at 10 mg/kg intraperitoneally on alternate days from Day –1 (1 day prior to collagen immunization) to Day 12. The animals were monitored clinically for the development of arthritis in the paws until Day 90. The apprint of arthritis was graded as: 0- normal; 1- arthritis localized to one joint; 2- more than one joint involved but not whole paw; 3- whole paw involved; 4- deformity of paw; 5- ankylosis of involved joints. The arthritis score of a mouse was the sum of the arthritic grades in each individual paw of the mouse (max = 20). The results are expressed as mean ± 5EM in each group.

The results, as shown in Figure 4, indicate that an arthritic score was measurable

in the C17.15-treated mice only after day 50 post-treatment, and that the peak mean arrhritic score obtained with the C17.15-treated mice was at least 5-fold lower than that measured in the IgG-treated mice. This demonstrated that the rat anti-mouse IL-12 antibody C17.15 prevented the development of collagen-induced arthritis in mice.

B. Suppression of Colitis in Mice by the Rat a-Murine IL-12 Antibody C17.15

IL-12 has also been demonstrated to play a role in the development/pathology of

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colitis. For example, anti-IL-12 antibodies have been shown to suppress disease in mouse models of colitis, e.g., TMBS induced colitis IL-2 knockout mice (Simpson et al. (1998) J. Exp. Med. 187(8): 1225-34). Similarly, anti-IL-2 antibodies have been demonstrated to suppress colitis formation in IL-10 knock-out mice. The ability of the rat anti-mouse IL-12 antibody, C17.15, to suppress TMBS colitis in mice was assessed in two studies (Davidson et al. (1998) J. Immunol. 161(6): 3143-9).

In the first study, colitis was induced in pathogen free SJL mice by the

administration of a 150 µL 50% ethanol solution containing 2.0 mg TNBS delivered via a pediatric umbilical artery catheter into the rectum. Control animals were treated with a 150 µL 50% ethanol solution only. A single dose of 0.75, 0.5, 0.5, 0.25, or 0.1 mg C17.15 or 0.75 mg control rat IgG2a was given intravenously via the tail vein at day 11, and the therapeutic effect of the treatment was assessed by weighing the animals on days 11 and therapeutic effect of the treatment was assessed by weighing the animals on days 11 and increased within 48 hours of antibody treatment and normalized on day 6 after treatment. The effect of treatment with C17.15 was confirmed histologically. Further,

assessments of IFN- γ secretion by CD4⁺ T-cells from spleen and colon of the treated mice, as well as IL-12 levels from spleen or colon-derived macrophages from the treated mice were also made (see Table 12).

In the second study, the dosing was optimized and the mice were treated with a

5 total dose of 0.1 mg or 0.5 mg C17.15 or 0.1 mg control IgG2a, respectively, split between days 12 and 14. It was found that the administration of C17.15 in a single dose at the dosage of 0.1 mg/mouse or 0.25 mg/mouse led to only partial improvement in TMBS-induced colitis and did not result in a significant reduction in the CD4⁺ T cell production of IFM-y in vitro, but did result in a significant decrease in secretion of IL-12, compared to untreated controls. At a single dose of 0.5 mg/mouse or greater a response was observed. Taking the lowest dose of antibody tested and administering it in two divided injections (at days 12 and 14) improved the dosing regimen, indicating that multiple low doses can be more effective than a single bolus dose. The data obtained are shown in Table 12.

Table 12: Anti-mouse II-12 mAb C17.15 Suppresses Established Colitis in Mice

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0	SEII	91.12	97.02		Ethanol control
	3489	86.71	16.2	gm 1.0 č1.71D	TABS + Ethanol
L	3618	<i>L. L</i> 1	16.28	gm 22.0 21.71D	TNBS + Ethanol
0	1723	⊅ 6′61	98.91	gm č.0 č1.71D	TNBS + Ethanol
0	7871	12.02	0.81	C17.15 0.75 mg	TABS + Ethanol
				gm č7.0	
300	3356	92.21	0.81	sadgl lonno	TNBS + Ethanol
		Day 17	Day 11		
(lm/gq)	(Jm/U)				
тасторнавеѕ	CD4, cells			Day 11	O ysa notion Day 0
IL-12 spieen	IEM-y spieen	ht (g)	giəW	Treatment	Disease

Administration of C17.15 monoclonal anti-IL-12 in two divided doses spaced

one day apart totaling 0.1 mg/mouse or 0.05 mg/mouse led to complete reversal of colitis as assessed by wasting and macroscopic appearance of the colon. In addition, this dose schedule led to significant down-regulation of lamina propria T-cell production of IFN- γ and macrophage production of IL-12, so that the latter were comparable to

levels seen in control ethanol-treated mice without TMBS-colitis. Thus, C17.15 administration to mouse models for TMBS colitis reversed the progression of the disease in a dose-dependent manner.

C. Suppression of Experimental Autoimmune Encephalomyelitis (EAE) in Mice by a-

IL-12 Antibodies

An a-IL-12 antibody was found to be able to inhibit the onset of acute EAE, to onset of acute EAE was assessed. by T cells of the TH₁ subset. Therefore, the ability of α -IL-12 antibodies to prevent the al. (1998) J. Exp. Med. 187: 537-546). The disease in this model is known to be induced 281-386; Banerjee, S. et al. (1998) Arthritis Rheum. (1998) 41: S33; and Segal, B.M. et encephalomyelitis (EAE), has been studied (Leonard, J.P. et al. (1995) J. Exp. Med. 181: pathogenesis of a murine model of multiple sclerosis, experimental autoimmune (Cormabella, M. et al. (1998) J. Clin. Invest. 102: 671-678). The role of IL-12 in the IL-12 production has been shown to correlate with disease activity in MS patients neurological diseases (Nicoletti, F. et al. (1996) J. Neuroimmunol. 70: 87-90). Increased 603). Elevated levels of serum IL-12 are detected in MS patients, but not in other antibodies in vitro (Balashov, K.E. et al. (1997) Proc. Natl. Acad. Sci. USA 94: 599-603). MS patients have enhanced IFN-y secretion that can be blocked with a-IL-12 CD40L expression (Balashov, K.E. et al. (1997) Proc. Natl. Acad. Sci. USA 94: 599-T cells) stimulate IL-12 production from antigen-presenting cells through unregulated A. et al. (1995) J. Exp. Med. 182: 1985-1996). T cells from MS patients (but not control acute plaques of MS patients but not in inflammatory brain infarct lesions (Windhagen, sclerosis (MS). The inducible IL-12 p40 message has been shown to be expressed in It is commonly believed that IL-12 plays a role in the pathogenesis of multiple

suppress the disease after onset, and to decrease the severity of relapses in mice immunized with the autoantigen, myelin basic protein (Banerjee, *S. et al.* (1998)

Arthritis Rheum. (1998) 41: S33). The beneficial effects of α-IL-12 antibody treatment in the mice persisted for over two months after stopping treatment. It has also been demonstrated that anti-IL-12 antibodies suppress the disease in mice that are recipients of encephalitogenic T cells by adoptive transfer (Leonard, J.P. et al. (1995) J. Exp. Med.

181: 281-386).

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EXAMPLE 9: Clinical Pharmacology of J695

In a double blind, crossover study, 64 healthy, human male subjects were administered ascending doses of J695 or placebo. Measurement of complement Ga prior to and 0.25 h after dosing did not demonstrate activation of the complement system. CRP and fibrinogen levels were only increased in subjects in whom symptoms of concurrent infections were observed.

All subjects survived and the overall tolerability of J695 was very good. In no

case did treatment have to be stopped because of adverse events (AEs). The most

o commonly observed AEs were headache and common cold/bronchitis, neither of which
were categorized as severe.

One of the study subjects, a 33-year-old single male, was suffering from psoriasis guttata at the start of the study. According to the randomized study design, this subject by chance received 5mg/kg 1695 by SC administration. Ten days prior to the arms and legs. At the time of the antibody administration, the subject displayed increased reddening, thickness of the erythematous plaques, and increased hyperkaratosis. One week after 1695 administration, the subject reported an hyperkaratosis. One week after 1695 administration, the lesions and a decrease in improvement in skin condition, including flattening of the lesions and a decrease in scaling. Shortly after the second administration of 1695 (5 mg/kg IV), the subject's skin scaling. Shortly after the second administration of 1695 (5 mg/kg IV), the subject's skin by scaling. Shortly after the second administration of 1695 (5 mg/kg IV), the subject's skin scaling. Shortly after the second administration of 1695 (5 mg/kg IV), the subject's skin by scaling. Shortly after the second administration of 1695 (5 mg/kg IV), the subject's skin by scaling covered of psoriatic lesions, in the absence of any local treatment.

25 EXAMPLE 10: Comparison of J695 Produced by Two CHO Cell Lines

expected clearance of J695 after the second administration of antibody.

For recombinant expression of J695, a recombinant expression vector encoding both the antibody heavy chain and the antibody light chain is introduced into dhfr- CHO cells (Urlaub, G. and Chasin, L.A. (1980) Proc. Natl. Acad. Sci. USA 77:4216-4220) by calcium phosphate-mediated transfection. Within the recombinant expression vector, the antibody heavy and light chain genes are each operatively linked to enhancer/promoter regulatory elements (e.g., derived from SV40, CMV, adenovirus and enhancer/promoter regulatory elements (e.g., derived from SV40, CMV, adenovirus and

the like, such as a CMV enhancer/AdMLP promoter regulatory element or an SV40 enhancer/AdMLP promoter regulatory element) to drive high levels of transcription of the genes. The recombinant expression vector also carries a DHFR gene, which allows for selection of CHO cells that have been transfected with the vector using methotrexate selection/amplification.

One hundred and fifty micrograms of an expression vector encoding the peptide sequences of the human antibody 1695 were dissolved in 2.7 ml water in a 50 ml conical tube. Three hundred µL of 2.5 M CaCl₂ were added and this DNA mixture was added dropwise to 3 ml of 2 x HEPES buffered saline in a 50 ml conical tube. After vortexing for 5 sec and incubating at room temperature for 20 min, 1 mL was distributed evenly over each plate (still in F12 medium), and the plates were incubated at 37 °C for evenly over each plate (still in F12 medium), and the plates were incubated at 37 °C for evenly over each plate (still in F12 medium), and the plates were incubated at 37 °C for evenly over each plate (still in F12 medium), and the plates were incubated at 37 °C for each plate. The DMSO shock continued for 1 min, after which the DMSO was diluted

 ${\rm CO}_2$ for two weeks, with one change of medium per week. Five days after the final medium change, culture supernatants were diluted 1:50

by the addition of 5 ml PBS to each plate. Plates were washed twice in PBS, followed by the addition of 10 ml of alpha MEM, supplemented with H/T and 5% FBS (selective for cells expressing DHFR) and overnight incubation at 37 °C. Cells were seeded into 96-well plates at a density of 100 cells per well, and plates were incubated at 37 °C, 5%

and tested using an ELISA specific for human IgG gamma chain. The clones yielding the highest ELISA signal were transferred from the 96-well plates to 12-well plates in 1.5 ml/well of Alpha MEM + 5% dialyzed serum. After 3 days, another ELISA specific for human IgG gamma chain was performed, and the 12 clones with the greatest activity were split into the alpha MEM + 5% dialyzed serum and 20 nM MTX. Cell line 031898 218 grew in the presence of 20 nM MTX without any apparent cell death or reduction in growth rate, produced 1.8 µg/ml hlgG in a three-day assay. T-25 cultures of 031898 218, growing in medium containing MTX, produced an average of 11.9 µg/ml of 1695. The line, designated ALP903, was adapted to growth in suspension under serum-free

medium, were passed again in 20 nM MTX. The cells were cultured under 100 nM MTX selection, followed by passaging in 500 nM MTX twice in the next 30 days. At

conditions, where it produced 7.5 pg J695/cell/24h.

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ALP903 cells, after initial selection in alpha MEM/5% FBS/20 nM MTX

that time the culture was producing 32 µg J695/mL/24 h. The culture was subcloned by limiting dilution. Subclone 218-22 produced 16.5 µg/mL in a 96-well plate in 2 days and 50.3 µg/mL of J695 in a 12-well dish in 2 days. Clone 218-22 was cultured in alpha MEM/5% dialyzed FBS/500 nM MTX for 38 days, followed by adaptation to serum-free free spinner culture, as above. The average cell-specific productivity of the serum-free suspension culture, designated ALP 905, was 58 pg/cell/24h.

The first cell line used to produce 1695 (ALP 903) resulted in lower yields of the antibody from culture than a second cell line, ALP 905. To assure that the ALP 905-produced 1695 was functionally identical to that produced from ALP 903, both batches of antibodies were assessed for IL-12 affinity, for the ability to block IL-12 binding to cellular receptors, for the ability to inhibit IFN-y induction by IL-12, and for the ability

to inhibit IL-12-mediated PHA blast proliferation.

The affinities of J695 batches ALP 903 and ALP 905 for IL-12 were determined

by measuring the kinetic rate constants of binding to IL-12 by surface plasmon resonance studies (BIAcore analyses). The off-rate constant (k_a) and the on-rate constant (k_a) of antibody batches ALP903 and ALP905 for binding to rhIL-12 were binding to IL-12 was calculated by dividing the off-rate constant by the on-rate constant. K_d was calculated for each separate experiment and then averaged. The results showed that the determined kinetic parameters and affinity of binding to rhIL-12 were very eightly for the determined kinetic parameters and affinity of binding to rhIL-12 were very eightly for the determined kinetic parameters and affinity of binding to rhIL-12 were very eightly for the for koff binding to the formula for the formula for the formula for the formula for the formula for the formula formula for the formula formula for the formula formula for formul

similar for J695 batches ALP 903 and ALP 905: the calculated K_d was 1.19 \pm 0.22 x 10° ¹⁰ M for batch ALP 905 (see Table 13). The ability of J695 derived from both ALP 903 and ALP 905 to block binding of

rhIL-12 to IL-12 receptors on human PHA-activated T-lymphoblasts was assessed (see Example 3). Each sample of 1695 was tested at a starting concentration of 1 x 10⁻⁸ with 10-fold serial dilutions. The antibody was preincubated for 1 hour at 37 °C with 50 pM [¹²⁵I]-human IL-12 in binding buffer. PHA blast cells were added to the antibody/[¹²⁵I]-human IL-12 mixture and incubated for 2 h at room temperature. Cell bound address the contraction of 1 x 10⁻⁸ with 30 pM and 1 x 10

radioactivity was separated from free [¹²⁵I]-IL-12 by centrifugation and washing steps, and % inhibition was calculated. The IC₅₀ values for J695 were determined from the inhibition curves using 4-parameter curve fitting and were confirmed by two

independent experiments. Incubations were carried out in duplicate. The results for the two batches of 1695 were very similar (see Table 13).

The ability of 1695 from both ALP 903 and ALP 905 cells to inhibit rhIL-12-

induced IFN- γ production by human PHA-activated lymphoblasts *in vitro* was assessed. Serial dilutions of 1695 were preincubated with 200 pg/mL rhIL-12 for 1 h at 37 °C. PHA lymphoblast cells were added and incubated for 18 hours at 37 °C. After incubation, cell free supernatant was withdrawn and the level of human IFN- γ determined by ELISA. The IC₅₀ values from the inhibition curves were plotted against the antibody concentration using 4-parameter curve fitting. The results demonstrate that

the ability of the two batches to inhibit IFN-y production is very similar.

The *in vitro* PHA blast cell proliferation assay was used to measure the neutralization capacity of ALP 903 and ALP 905 1695 for rhIL-12. Serial dilutions of 1695 of each type were preincubated with 230 pg/mL human IL-12 for 1 h at 37 °C. Mext PHA blast cells were added and incubated for 3 days at 37 °C. The cells were then labeled for 6 hours with 1 yCi/well [³H]-thymidine. The cultures were harvested and labeled for 6 hours with 1 yCi/well [³H]-thymidine incorporation measured. Non-specific proliferation (background) was

[³H]-thymidine incorporation measured. Non-specific proliferation (background) was measured in the absence of rhIL-12. The IC₅₀ values for ALP 903 and ALP 905 1695 were found to be very similar and are set forth in Table 13.

The activity of the J695 antibodies in neutralizing rhIL-12 activity, in blocking

20 IL-12 binding to cell surface receptors, and in binding to rhIL-12 did not significantly differ from batch ALP 903 to batch ALP 905, and thus the antibodies produced from these two different cell types were equivalent.

Table 13: Comparison of the Properties of 1695 lots ALP 903 and ALP 905

						¥LP 905
4.3 x 10 ⁻¹²	4.4 x 10 ⁻¹²	11.01 x 0.8	⁰¹⁻ 01 x 94.1	^{c-} 01 x 62.2	3.91 x 10 ⁵	5691
						¥Fb 903
²¹ ·01 x 8.č	²¹⁻ 01 x č.č	71-01 x 4.8	01-01 × 61.1	c01×9++	3.75 × 10 ⁵	5691
(M)	(M)					
Assay IC ₅₀	Assay IC ₅₀	IC ²⁰ (M)		(₁ .s)	('R'', S')	
IEN-1	teald AH9	KB assay	$K^q(M)$	k _d , Off-rate	k _a , On-rate	*Kpodiju

EQUIVALENTS

Those skilled in the art will recognize, or be able to ascertain using no more than routine experimentation, many equivalents to the specific embodiments of the invention described herein. Such equivalents are intended to be encompassed by the following

claims

CLAIMS

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- I. An isolated human antibody, or an antigen-binding portion thereof, that binds to human IL-12, wherein said human antibody is a neutralizing antibody.
- 2. A selectively mutated human IL-12 antibody, comprising: a human antibody or antigen-binding portion thereof, selectively mutated at a preferred selective mutagenesis, contact or hypermutation position with an activity enhancing amino acid residue such that it binds to human IL-12.
- 3. A selectively mutated human IL-12 antibody, comprising: a human antibody or antigen-binding portion thereof, selectively mutated at a preferred selective mutagenesis position with an activity enhancing amino acid residue such that it binds to human IL-12.
- 4. The selectively mutated human IL-12 antibody of claim 2, wherein the human antibody or antigen-binding portion thereof is selectively mutated at more than one preferred selective mutagenesis, contact or hypermutation positions with an activity enhancing amino acid residue.
- 5. The selectively mutated human IL-12 antibody of claim 4, wherein the human antibody or antigen-binding portion thereof is selectively mutated at no more than three preferred selective mutagenesis, contact or hypermutation positions.
- The selectively mutated human IL-12 antibody of claim 4, wherein the human antibody or antigen-binding portion thereof is selectively mutated at no more than two preferred selective mutagenesis, contact or hypermutation positions.

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7. The selectively mutated human IL-12 antibody of claim 2, wherein the human antibody or antigen-binding portion thereof, is selectively mutated such that a target specificity affinity level is attained, said target level being improved over that attainable when selecting for an antibody against the same antigen using phage display technology.

8. The selectively mutated human IL-12 antibody of claim 7, wherein the selectively mutated human antibody further retains at least one desirable property or characteristic.

- 9. An isolated human antibody, or antigen-binding portion thereof, that binds to human IL-12 and dissociates from human IL-12 with a k_{off} rate constant of 0.1s⁻¹ or less, as determined by surface plasmon resonance, or which inhibits phytohemagglutinin blast proliferation in an *in vitro* phytohemagglutinin blast proliferation assay (PHA assay) with an IC₅₀ of 1 x 10⁻⁶M or less.
- 10. The isolated human antibody of claim 9, or an antigen-binding portion thereof, which dissociates from human IL-12 with a k_{off} rate constant of 1 x 10^{-2} s⁻¹or less, as determined by surface plasmon resonance, or which inhibits phytohemagglutinin blast proliferation in an *in vitro* PHA assay with an IC₅₀ of 1 x 10^{-7} M or less.
- 11. The isolated human antibody of claim 9, or an antigen-binding portion thereof, which dissociates from human IL-12 with a $k_{\rm off}$ rate constant of 1 x 10⁻³s⁻¹ or less, as determined by surface plasmon resonance, or which inhibits phytohemagglutinin blast proliferation in an *in vitro* PHA assay with an IC₅₀ of 1 x 10⁻⁸M or less.
- 12. The isolated human antibody of claim 9, or an antigen-binding portion thereof, which dissociates from human IL-12 with a $k_{\rm off}$ rate constant of 1 x $10^{-4} s^{-1}$ or less, as determined by surface plasmon resonance, or which inhibits phytohemagglutinin blast proliferation in an *in vitro* PHA assay with an IC₅₀ of 1 x 10^{-9} M or less.

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13. The isolated human antibody of claim 9, or an antigen-binding portion thereof, which dissociates from human IL-12 with a $k_{\rm off}$ rate constant of 1 x 10^{-5} s⁻¹ or less, as determined by surface plasmon resonance, or which inhibits phytohemagglutinin blast proliferation in an *in vitro* PHA assay with an IC₅₀ of 1 x 10^{-10} M or less.

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14. The isolated human antibody of claim 9, or an antigen-binding portion thereof, which dissociates from human IL-12 with a $k_{\rm off}$ rate constant of 1 x 10^{-5} s⁻¹ or less, as determined by surface plasmon resonance, or which inhibits phytohemagglutinin blast proliferation in an *in vitro* PHA assay with an IC₅₀ of 1 x 10^{-11} M or less.

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- 15. An isolated human antibody, or an antigen-binding portion thereof, which has the following characteristics:
- a) inhibits phytohemagglutinin blast proliferation in an *in vitro* PHA assay with an IC₅₀ of 1 x 10^{-6} M or less;
- b) has a heavy chain CDR3 comprising the amino acid sequence of SEQ ID NO: 1; and
- c) has a light chain CDR3 comprising the amino acid sequence of SEQ ID NO: 2.
- 16. The isolated human antibody of claim 15, or an antigen-binding portion thereof, which further has a heavy chain CDR2 comprising the amino acid sequence of SEQ ID NO: 3; and has a light chain CDR2 comprising the amino acid sequence of SEQ ID NO: 4.
- 17. The isolated human antibody of claim 15, or an antigen-binding portion thereof, which further has a heavy chain CDR1 comprising the amino acid sequence of SEQ ID NO: 5; and has a light chain CDR1 comprising the amino acid sequence of SEQ ID NO: 6.

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18. The isolated human antibody, or antigen binding portion thereof of claim 16, which has a heavy chain variable region comprising the amino acid sequence of SEQ ID NO: 7; and has a light chain variable region comprising the amino acid sequence of SEQ ID NO: 8.

- 19. An isolated human antibody, or an antigen-binding portion thereof, which has the following characteristics:
- a) inhibits phytohemagglutinin blast proliferation in an *in vitro* PHA assay with an IC_{50} of 1 x 10^{-9} M or less;
- b) has a heavy chain CDR3 comprising the amino acid sequence of SEQ IDNO: 9; and
 - c) has a light chain CDR3 comprising the amino acid sequence of SEQ ID
 NO: 10.
- 15 20. The isolated human antibody of claim 19, or an antigen-binding portion thereof, which further has a heavy chain CDR2 comprising the amino acid sequence of SEQ ID NO: 11; and has a light chain CDR2 comprising the amino acid sequence of SEQ ID NO: 12.
- 21. The isolated human antibody of claim 19, or an antigen-binding portion thereof, which further has a heavy chain CDR1 comprising the amino acid sequence of SEQ ID NO: 13; and has a light chain CDR1 comprising the amino acid sequence of SEQ ID NO: 14.
- 25. The isolated human antibody of claim 19, which has a heavy chain variable region comprising the amino acid sequence of SEQ ID NO: 15; and has a light chain variable region comprising the amino acid sequence of SEQ ID NO: 16.

- 23. An isolated human antibody, or an antigen-binding portion thereof, which
- a) inhibits phytohemagglutinin blast proliferation in an *in vitro* PHA assay with an IC₅₀ of 1 x 10^{-9} M or less;
- b) has a heavy chain CDR3 comprising the amino acid sequence of SEQ ID NO: 17; and
 - c) has a light chain CDR3 comprising the amino acid sequence of SEQ ID NO: 18.
- 24. The isolated human antibody, or an antigen-binding portion thereof, of claim 23 which further has a heavy chain CDR2 comprising the amino acid sequence of SEQ ID NO: 19; and a light chain CDR2 comprising the amino acid sequence of SEQ ID NO: 20.
- The isolated human antibody, or an antigen-binding portion thereof, of
 claim 23 which further has a heavy chain CDR1 comprising the amino acid sequence of
 SEQ ID NO: 21; and a light chain CDR1 comprising the amino acid sequence of SEQ
 ID NO: 22.
- 26. An isolated human antibody, or an antigen-binding portion thereof,
 having a heavy chain variable region comprising the amino acid sequence of SEQ ID
 NO: 23, and a light chain variable region comprising the amino acid sequence of SEQ
 ID NO: 24.
- 27. The isolated human antibody of claim 26, comprising a heavy chain constant region selected from the group consisting of IgG1, IgG2, IgG3, IgG4, IgM, IgA and IgE constant regions.
 - 28. The isolated human antibody of claim 27, wherein the antibody heavy chain constant region is IgG1.

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29. The isolated human antibody of claim 26, which is a Fab fragment.

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- 30. The isolated human antibody of claim 26, which is a F(ab')₂ fragment.
- 31. The isolated human antibody of claim 26, which is a single chain Fv fragment.

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- 32. An isolated human antibody, or an antigen-binding portion thereof, which
- a) inhibits phytohemagglutinin blast proliferation in an *in vitro* PHA assay with an IC₅₀ of 1 x 10^{-9} M or less;
- b) has a heavy chain CDR3 comprising the amino acid sequence selected from the group consisting of SEQ ID NO: 404-SEQ ID NO: 469; or
 - c) has a light chain CDR3 comprising the amino acid sequence selected from the group consisting of SEQ ID NO: 534-SEQ ID NO: 579.
- 33. The isolated human antibody, or an antigen-binding portion thereof, of claim 32 which further has a heavy chain CDR2 comprising the amino acid sequence selected from the group consisting of SEQ ID NO:335-SEQ ID NO: 403; or a light chain CDR2 comprising the amino acid sequence selected from the group consisting of SEQ ID NO: 506-SEQ ID NO: 533.
- 34. The isolated human antibody, or an antigen-binding portion thereof, of claim 32 which further has a heavy chain CDR1 comprising the amino acid sequence selected from the group consisting of SEQ ID NO: 288-SEQ ID NO: 334; or a light chain CDR1 comprising the amino acid sequence selected from the group consisting of SEQ ID NO: 470-SEQ ID NO: 505.

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35. An isolated human antibody, or an antigen-binding portion thereof, having a the heavy chain variable region comprising the amino acid sequence of SEQ ID NO: 23, and a light chain variable region comprising the amino acid sequence of SEQ ID NO: 24.

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- 36. The isolated human antibody of claim 35, comprising a heavy chain constant region selected from the group consisting of IgG1, IgG2, IgG3, IgG4, IgM, IgA and IgE constant regions.
- 5 37. The isolated human antibody of claim 36, wherein the antibody heavy chain constant region is IgG1.
 - 38. The isolated human antibody of claim 35, which is a Fab fragment.
- The isolated human antibody of claim 35, which is a F(ab')? fragment.
 - 40. The isolated human antibody of claim 35, which is a single chain Fv fragment.
- 15 41. An isolated human antibody, or an antigen-binding portion thereof, which
 a) inhibits phytohemagglutinin blast proliferation in an *in vitro* PHA assay with an IC₅₀ of 1 x 10⁻⁹M or less;
 - b) has a heavy chain CDR3 comprising the amino acid sequence of SEQ ID NO: 25; and
- 20 c) has a light chain CDR3 comprising the amino acid sequence of SEQ ID NO: 26.
 - 42. The isolated human antibody, or an antigen-binding portion thereof, of claim 41 which further has a heavy chain CDR2 comprising the amino acid sequence of SEQ ID NO: 27; and a light chain CDR2 comprising the amino acid sequence of SEQ ID NO: 28.
 - 43. The isolated human antibody, or an antigen-binding portion thereof, of claim 41 which further has a heavy chain CDR1 comprising the amino acid sequence of SEQ ID NO: 29; and a light chain CDR1 comprising the amino acid sequence of SEQ ID NO: 30.

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44. An isolated human antibody, or an antigen-binding portion thereof, having a heavy chain variable region comprising the amino acid sequence of SEQ ID NO: 31, and a light chain variable region comprising the amino acid sequence of SEQ ID NO: 32.

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- 45. The isolated human antibody of claim 44, comprising a heavy chain constant region selected from the group consisting of IgG1, IgG2, IgG3, IgG4, IgM, IgA and IgE constant regions.
- 10 46. The isolated human antibody of claim 45, wherein the antibody heavy chain constant region is IgG1.
 - 47. The isolated human antibody of claim 44, which is a Fab fragment.
- The isolated human antibody of claim 44, which is a F(ab')₂ fragment.
 - 49. The isolated human antibody of claim 44, which is a single chain Fv fragment.
 - 50. An isolated human antibody, or an antigen-binding portion thereof, which
 - a) inhibits phytohemagglutinin blast proliferation in an *in vitro* PHA assay with an IC₅₀ of 1 x 10^{-6} M or less;
 - b) comprises a heavy chain CDR3 comprising the amino acid sequence of SEQ ID NO: 1, a heavy chain CDR2 comprising the amino acid sequence of SEQ ID NO: 3 and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID NO: 5, or a mutant thereof having one or more amino acid substitutions at a contact position or a hypermutation position, wherein said mutant has a k_{off} rate no more than 10-fold higher than the antibody comprising a heavy chain CDR3 comprising the amino acid sequence of SEQ ID NO: 1, a heavy chain CDR2 comprising the amino acid sequence of SEQ ID NO: 3, and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID NO: 5; and

- c) comprises a light chain CDR3 comprising the amino acid sequence of SEQ ID NO: 2, a light chain CDR2 comprising the amino acid sequence of SEQ ID NO: 4, and a light chain CDR1 comprising the amino acid sequence of SEQ ID NO: 6, or a mutant thereof having one or more amino acid substitutions at a contact position or a hypermutation position, wherein said mutant has a k_{off} rate no more than 10-fold higher than the antibody comprising a light chain CDR3 comprising the amino acid sequence of SEQ ID NO: 2, a light chain CDR2 comprising the amino acid sequence of SEQ ID NO: 4, and a light chain CDR1 comprising the amino acid sequence of SEQ ID NO: 6.
- 10 51. An isolated human antibody, or an antigen-binding portion thereof, which
 - a) inhibits phytohemagglutinin blast proliferation in an *in vitro* PHA assay with an IC₅₀ of 1 x 10^{-9} M or less;
 - b) comprises a heavy chain CDR3 comprising the amino acid sequence of SEQ ID NO: 9, a heavy chain CDR2 comprising the amino acid sequence of SEQ ID NO: 11 and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID NO: 13, or a mutant thereof having one or more amino acid substitutions at a contact position or a hypermutation position, wherein said mutant has a k_{off} rate no more than 10-fold higher than the antibody comprising a heavy chain CDR3 comprising the amino acid sequence of SEQ ID NO: 9, a heavy chain CDR2 comprising the amino acid sequence of SEQ ID NO: 11, and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID NO: 13; and
 - c) comprises a light chain CDR3 comprising the amino acid sequence of SEQ ID NO: 10, a light chain CDR2 comprising the amino acid sequence of SEQ ID NO: 12, and a light chain CDR1 comprising the amino acid sequence of SEQ ID NO: 14, or a mutant thereof having one or more amino acid substitutions at a contact position or a hypermutation position, wherein said mutant has a k_{off} rate no more than 10-fold higher than the antibody comprising a light chain CDR3 comprising the amino acid sequence of SEQ ID NO: 10, a light chain CDR2 comprising the amino acid sequence of SEQ ID NO: 12, and a light chain CDR1 comprising the amino acid sequence of SEQ ID NO: 14.

- 52. An isolated human antibody, or an antigen-binding portion thereof, which
- a) inhibits phytohemagglutinin blast proliferation in an *in vitro* PHA assay with an IC₅₀ of 1 x 10^{-9} M or less;
- b) comprises a heavy chain CDR3 comprising the amino acid sequence of SEQ ID NO: 17, a heavy chain CDR2 comprising the amino acid sequence of SEQ ID NO: 19 and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID NO: 21, or a mutant thereof having one or more amino acid substitutions at a contact position or a hypermutation position, wherein said mutant has a k_{off} rate no more than 10-fold higher than the antibody comprising a heavy chain CDR3 comprising the amino acid sequence of SEQ ID NO: 17, a heavy chain CDR2 comprising the amino acid sequence of SEQ ID NO: 19, and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID NO: 21; and
 - c) comprises a light chain CDR3 comprising the amino acid sequence of SEQ ID NO: 18, a light chain CDR2 comprising the amino acid sequence of SEQ ID NO: 20, and a light chain CDR1 comprising the amino acid sequence of SEQ ID NO: 22, or a mutant thereof having one or more amino acid substitutions at a contact position or a hypermutation position, wherein said mutant has a k_{off} rate no more than 10-fold higher than the antibody comprising a light chain CDR3 comprising the amino acid sequence of SEQ ID NO: 18, a light chain CDR2 comprising the amino acid sequence of SEQ ID NO: 20, and a light chain CDR1 comprising the amino acid sequence of SEQ ID NO: 22.
 - 53. An isolated nucleic acid encoding the heavy chain CDR3 comprising the amino acid sequence of SEQ ID NO: 17.
 - 54. The isolated nucleic acid of claim 53, which encodes an antibody heavy chain variable region.

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55. The isolated nucleic acid of claim 54, wherein the CDR2 of the antibody heavy chain variable region comprises the amino acid sequence of SEQ ID NO: 19.

- 56. The isolated nucleic acid of claim 54, wherein the CDR1 of the antibody heavy chain variable region comprises the amino acid sequence of SEQ ID NO: 21.
- 57. The isolated nucleic acid of claim 56, which encodes an antibody heavy chain variable region comprising the amino acid sequence of SEQ ID NO: 23.
 - 58. An isolated nucleic acid encoding the light chain CDR3 comprising the amino acid sequence of SEQ ID NO: 18.
- 10 59. The isolated nucleic acid of claim 58, which encodes an antibody light chain variable region.
 - 60. The isolated nucleic acid of claim 59, wherein the CDR2 of the antibody light chain variable region comprises the amino acid sequence of SEQ ID NO: 20.
 - 61. The isolated nucleic acid of claim 59, wherein the CDR1 of the antibody light chain variable region comprises the amino acid sequence of SEQ ID NO: 22.
- 62. The isolated nucleic acid of claim 61, which encodes an antibody light chain variable region comprising the amino acid sequence of SEQ ID NO: 24.

- 63. An isolated human antibody, or an antigen-binding portion thereof, which
- a) inhibits phytohemagglutinin blast proliferation in an *in vitro* PHA assay with an IC_{50} of 1 x 10^{-9} M or less;
- b) comprises a heavy chain CDR3 comprising the amino acid sequence of SEQ ID NO: 25, a heavy chain CDR2 comprising the amino acid sequence of SEQ ID NO: 27 and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID NO: 29, or a mutant thereof having one or more amino acid substitutions at a contact position or a hypermutation position, wherein said mutant has a k_{off} rate no more than 10-fold higher than the antibody comprising a heavy chain CDR3 comprising the amino acid sequence of SEQ ID NO: 25, a heavy chain CDR2 comprising the amino acid sequence

of SEQ ID NO: 27, and a heavy chain CDR1 comprising the amino acid sequence of SEQ ID NO: 29; and

- c) comprises a light chain CDR3 comprising the amino acid sequence of SEQ ID NO: 26, a light chain CDR2 comprising the amino acid sequence of SEQ ID NO: 28, and a light chain CDR1 comprising the amino acid sequence of SEQ ID NO: 30, or a mutant thereof having one or more amino acid substitutions at a contact position or a hypermutation position, wherein said mutant has a k_{off} rate no more than 10-fold higher than the antibody comprising a light chain CDR3 comprising the amino acid sequence of SEQ ID NO: 26, a light chain CDR2 comprising the amino acid sequence of SEQ ID NO: 28, and a light chain CDR1 comprising the amino acid sequence of SEQ ID NO: 30.
 - 64. An isolated nucleic acid encoding the heavy chain CDR3 comprising the amino acid sequence of SEQ ID NO: 25.

65. The isolated nucleic acid of claim 64, which encodes an antibody heavy chain variable region.

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- 66. The isolated nucleic acid of claim 65, wherein the CDR2 of the antibody heavy chain variable region comprises the amino acid sequence of SEQ ID NO: 27.
 - 67. The isolated nucleic acid of claim 65, wherein the CDR1 of the antibody heavy chain variable region comprises the amino acid sequence of SEQ ID NO: 29.
- 68. The isolated nucleic acid of claim 67, which encodes an antibody heavy chain variable region comprising the amino acid sequence of SEQ ID NO: 31.
 - 69. An isolated nucleic acid encoding the light chain CDR3 comprising the amino acid sequence of SEQ ID NO: 26.
 - 70. The isolated nucleic acid of claim 69, which encodes an antibody light chain variable region.

- 71. The isolated nucleic acid of claim 70, wherein the CDR2 of the antibody light chain variable region comprises the amino acid sequence of SEQ ID NO: 28.
- 5 72. The isolated nucleic acid of claim 70, wherein the CDR1 of the antibody light chain variable region comprises the amino acid sequence of SEQ ID NO: 30.
 - 73. The isolated nucleic acid of claim 72, which encodes an antibody heavy chain variable region comprising the amino acid sequence of SEQ ID NO: 32.

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- 74. An isolated human antibody, or an antigen-binding portion thereof, which has the following characteristics:
- a) that binds to human IL-12 and dissociates from human IL-12 with a $k_{\rm off}$ rate constant of 0.1 s⁻¹ or less, as determined by surface plasmon resonance, or which inhibits phytohemagglutinin blast proliferation in an *in vitro* phytohemagglutinin blast proliferation assay (PHA assay) with an IC₅₀ of 1 x 10⁻⁶M or less.
- b) has a heavy chain variable region comprising an amino acid sequence selected from a member of the VH3 germline family, wherein the heavy chain variable region has a mutation at a contact or hypermutation position with an activity enhancing amino acid residue.
- c) has a light chain variable region comprising an amino acid sequence selected from a member of the $V\lambda 1$ germline family, wherein the light chain variable region has a mutation at a contact or hypermutation position with an activity enhancing amino acid residue.

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- 75. An isolated human antibody, or an antigen-binding portion thereof, which has the following characteristics:
- a) that binds to human IL-12 and dissociates from human IL-12 with a $k_{\rm off}$ rate constant of $0.1s^{-1}$ or less, as determined by surface plasmon resonance, or which inhibits phytohemagglutinin blast proliferation in an *in vitro* phytohemagglutinin blast proliferation assay (PHA assay) with an IC₅₀ of 1 x 10⁻⁶M or less.
- b) has a heavy chain variable region comprising an amino acid sequence selected from the group consisting of SEQ ID NOs: 595-667, wherein the heavy chain variable region has a mutation at a contact or hypermutation position with an activity enhancing amino acid residue.
- c) has a light chain variable region comprising an amino acid sequence selected from the group consisting of SEQ ID NOs: 669-675, wherein the light chain variable region has a mutation at a contact or hypermutation position with an activity enhancing amino acid residue.

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- 76. An isolated human antibody, or an antigen-binding portion thereof, which has the following characteristics:
- a) that binds to human IL-12 and dissociates from human IL-12 with a $k_{\rm off}$ rate constant of 0.1 s⁻¹or less, as determined by surface plasmon resonance, or which inhibits phytohemagglutinin blast proliferation in an *in vitro* phytohemagglutinin blast proliferation assay (PHA assay) with an IC₅₀ of 1 x 10⁻⁶M or less.
- b) has a heavy chain variable region comprising the COS-3 germline amino acid sequence, wherein the heavy chain variable region has a mutation at a contact or hypermutation position with an activity enhancing amino acid residue.
- c) has a light chain variable region comprising the DPL8 germline amino acid sequence, wherein the light chain variable region has a mutation at a contact or hypermutation position with an activity enhancing amino acid residue.

77. An isolated human antibody, or an antigen-binding portion thereof, which has the following characteristics:

- a) that binds to human IL-12 and dissociates from human IL-12 with a $k_{\rm off}$ rate constant of 0.1 s⁻¹or less, as determined by surface plasmon resonance, or which inhibits phytohemagglutinin blast proliferation in an *in vitro* phytohemagglutinin blast proliferation assay (PHA assay) with an IC₅₀ of 1 x 10⁻⁶M or less.
- b) has a heavy chain variable region comprising an amino acid sequence selected from a member of the VH3 germline family, wherein the heavy chain variable region comprises a CDR2 that is structurally similar to CDR2s from other VH3 germline family members, and a CDR1 that is structurally similar to CDR1s from other VH3 germline family members, and wherein the heavy chain variable region has a mutation at a contact or hypermutation position with an activity enhancing amino acid residue;
- c) has a light chain variable region comprising an amino acid sequence selected from a member of the V λ 1 germline family, wherein the light chain variable region comprises a CDR2 that is structurally similar to CDR2s from other V λ 1 germline family members, and a CDR1 that is structurally similar to CDR1s from other V λ 1 germline family members, and wherein the light chain variable region has a mutation at a contact or hypermutation position with an activity enhancing amino acid residue.
- The isolated human antibody, or antigen binding portion thereof, of claims 74, 75, 76, or 77, wherein the mutation is in the heavy chain CDR3.
 - 79. The isolated human antibody, or antigen binding portion thereof, of claims 74, 75, 76, or 77, wherein the mutation is in the light chain CDR3.
 - 80. The isolated human antibody, or antigen binding portion thereof, of claims 74, 75, 76, or 77, wherein the mutation is in the heavy chain CDR2.

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81. The isolated human antibody, or antigen binding portion thereof, of claims 74, 75, 76, or 77, wherein the mutation is in the light chain CDR2.

- 82. The isolated human antibody, or antigen binding portion thereof, of claims 74, 75, 76, or 77, wherein the mutation is in the heavy chain CDR1.
- 83. The isolated human antibody, or antigen binding portion thereof, of claims 74, 75, 76, or 77, wherein the mutation is in the light chain CDR1.
 - 84. A recombinant expression vector encoding:
 - a) an antibody heavy chain having a variable region comprising the amino acid sequence of SEQ ID NO: 31; and
- b) an antibody light chain having a variable region comprising the amino acid sequence of SEQ ID NO: 32.
 - 85. A host cell into which the recombinant expression vector of claim 84 has been introduced.
 - 86. A method of synthesizing a human antibody that binds human IL-12, comprising culturing the host cell of claim 85 in a culture medium until a human antibody that binds human IL-12 is synthesized by the cell.
- 20 87. An isolated human antibody, or antigen-binding portion thereof, that neutralizes the activity of human IL-12, and at least one additional primate IL-12 selected from the group consisting of baboon IL-12, marmoset IL-12, chimpanzee IL-12, cynomolgus IL-12 and rhesus IL-12, but which does not neutralize the activity of the mouse IL-12.
 - 88. A pharmaceutical composition comprising the antibody or an antigen binding portion thereof, of any one of claims 1-52, 74-83 and 87 and a pharmaceutically acceptable carrier.
- 30 89. A composition comprising the antibody or an antigen binding position thereof, of any one of claims 1-52, 74-83 and 87 and an additional agent.

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- 90. The composition of claim 89, wherein the additional agent is a therapeutic agent.
- 91. The composition of claim 90, wherein the therapeutic agent is selected from the group consisting of budenoside, epidermal growth factor, corticosteroids, cyclosporin, sulfasalazine, aminosalicylates, 6-mercaptopurine, azathioprine, metronidazole, lipoxygenase inhibitors, mesalamine, olsalazine, balsalazide, antioxidants, thromboxane inhibitors. IL-1 receptor antagonists, anti-IL-1β monoclonal antibodies, anti-IL-6 monoclonal antibodies, growth factors, elastase inhibitors, pyridinyl-imidazole compounds, antibodies or agonists of TNF, LT, IL-1, IL-2, IL-6, IL-7, IL-8, IL-15, IL-16, IL-18, EMAP-II, GM-CSF, FGF, and PDGF, antibodies of CD2, CD3, CD4, CD8, CD25, CD28, CD30, CD40, CD45, CD69, CD90 or their ligands, methotrexate, cyclosporin, FK506, rapamycin, mycophenolate mofetil, leflunomide, NSAIDs, ibuprofen, corticosteroids, prednisolone, phosphodiesterase inhibitors, adenosine agonists, antithrombotic agents, complement inhibitors, adrenergic agents, IRAK, NIK, IKK, p38, MAP kinase inhibitors, IL-1β converting enzyme inhibitors, TNFα converting enzyme inhibitors, T-cell signalling inhibitors, metalloproteinase inhibitors, sulfasalazine, azathioprine, 6-mercaptopurines, angiotensin converting enzyme inhibitors, soluble cytokine receptors, soluble p55 TNF receptor, soluble p75 TNF receptor, sIL-1RI, sIL-1RII, sIL-6R, antiinflammatory cytokines, IL-4, IL-10, IL-11, IL-13 and TGFβ.
 - 92. The therapeutic composition of claim 90, wherein the therapeutic agent is selected from the group consisting of anti-TNF antibodies and antibody fragments thereof, TNFR-Ig constructs, TACE inhibitors, PDE4 inhibitors, corticosteroids, budenoside, dexamethasone, sulfasalazine, 5-aminosalicylic acid, olsalazine, IL-1β converting enzyme inhibitors, IL-1ra, tyrosine kinase inhibitors, 6-mercaptopurines and IL-11.
 - 93. The therapeutic composition of claim 90, wherein the therapeutic agent is selected from the group consisting of corticosteroids, prednisolone, methylprednisolone, azathioprine, cyclophosphamide, cyclosporine, methotrexate, 4-aminopyridine,

tizanidine, interferon-β1a, interferon-β1b. Copolymer 1, hyperbaric oxygen, intravenous immunoglobulin, clabribine, antibodies or agonists of TNF, LT, IL-1, IL-2, IL-6, IL-7, IL-8, IL-15, IL-16, IL-18, EMAP-II. GM-CSF, FGF. PDGF, antibodies to CD2, CD3, CD4, CD8, CD25, CD28, CD30, CD40, CD45, CD69, CD80, CD86, CD90 or their ligands, methotrexate, cyclosporine, FK506, rapamycin, mycophenolate mofetil, leflunomide, NSAIDs, ibuprofen, corticosteroids, prednisolone, phosphodiesterase inhibitors, adensosine agonists, antithrombotic agents, complement inhibitors, adrenergic agents, IRAK, NIK, IKK, p38 or MAP kinase inhibitors, IL-1β converting enzyme inhibitors, T-cell signalling inhibitors, kinase inhibitors, metalloproteinase inhibitors, sulfasalazine, azathioprine, 6-mercaptopurines, angiotensin converting enzyme inhibitors, soluble cytokine receptors, soluble p55 TNF receptor, soluble p75 TNF receptor, sIL-1RI, sIL-1RII, sIL-6R, sIL-13R, anti-P7s, p-selectin glycoprotein ligand (PSGL), antiinflammatory cytokines, IL-4, IL-10, IL-13 and TGFβ.

94. A method for inhibiting human IL-12 activity comprising contacting human IL-12 with the antibody of claim 44 such that human IL-12 activity is inhibited.

- 95. A method for inhibiting human IL-12 activity in a human subject suffering from a disorder in which IL-12 activity is detrimental, comprising administering to the human subject the antibody of claim 44 such that human IL-12 activity in the human subject is inhibited.
- 96. The method of claim 95, wherein the disorder is selected from the group consisting of rheumatoid arthritis, osteoarthritis, juvenile chronic arthritis,

 Lyme arthritis, psoriatic arthritis, reactive arthritis, spondyoarthropathy, ankylosing spondylitis, systemic lupus erythematosis, Crohn's disease, ulcerative colitis, inflammatory bowel disease, multiple sclerosis, insulin dependent diabetes mellitus, thyroiditis, asthma, allergic diseases, psoriasis, dermatitisscleroderma, thyroiditis, graft versus host disease, organ transplant rejection, acute or chronic immune disease associated with organ transplantation, sarcoidosis, atherosclerosis, disseminated intravascular coagulation, Kawasaki's disease, Grave's disease, nephrotic syndrome, chronic fatigue syndrome, polyarteritis nodosa, Wegener's granulomatosis, Henoch-

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Schonlein purpura, microscopic vasculitis of the kidneys, chronic active hepatitis, Sjogren's syndrome, uvcitis, sepsis, septic shock, sepsis syndrome, adult respiratory distress syndrome, cachexia, infectious diseases, parasitic diseases, acquired immunodeficiency syndrome, acute transverse myelitis, myasthenia gravis, Huntington's chorea, Parkinson's disease, Alzheimer's disease, stroke, primary biliary cirrhosis, fibrotic lung diseases, hemolytic anemia, malignancies, heart failure and myocardial infarction.

- 97. The method of claim 95, wherein the disorder is Crohn's disease.
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- 98. The method of claim 95, wherein the disorder is multiple sclerosis.
- 99. The method of claim 95, wherein the disorder is rheumatoid arthritis.
- 15 100. A method for improving the activity of an antibody, or antigen-binding portion thereof, to attain a predetermined target activity, comprising:
 - a) providing a parent antibody a antigen-binding portion thereof;
 - b) selecting a preferred selective mutagenesis position selected from group consisting of H30, H31, H31B, H32, H33, H52, H56, H58, L30, L31, L32, L50, L91, L92, L93, L94.
 - c) individually mutating the selected preferred selective mutagenesis position to at least two other amino acid residues to hereby create a first panel of mutated antibodies, or antigen binding portions thereof;
- d) evaluating the activity of the first panel of mutated antibodies, or
 antigen binding portions thereof to determined if mutation of a single selective
 mutagenesis position produces an antibody or antigen binding portion thereof with the
 predetermined target activity or a partial target activity;
 - e) combining in a stepwise fashion, in the parent antibody, or antigen binding portion thereof, individual mutations shown to have an improved activity, to form combination antibodies, or antigen binding portions thereof.

- f) evaluating the activity of the combination antibodies, or antigen binding portions thereof to determined if the combination antibodies, or antigen binding portions thereof have the predetermined target activity or a partial target activity.
- g) if steps d) or f) do not result in an antibody or antigen binding portion thereof having the predetermined target activity, or result an antibody with only a partial activity, the method further comprising mutating additional amino acid residues selected from the group consisting of H35, H50, H53, H54, H95, H96, H97, H98, L30A and L96 to at least two other amino acid residues to thereby create a second panel of mutated antibodies or antigen-binding portions thereof;
 - h) evaluating the activity of the second panel of mutated antibodies or antigen binding portions thereof, to determined if mutation of a single amino acid residue selected from the group consisting of H35, H50, H53, H54, H95, H96, H97, H98, L30A and L96 results an antibody or antigen binding portion thereof, having the predetermined target activity or a partial activity;

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- i) combining in stepwise fashion in the parent antibody, or antigenbinding portion thereof, individual mutations of step g) shown to have an improved activity, to form combination antibodies, or antigen binding portions thereof;
- j) evaluating the activity of the combination antibodies or antigen binding portions thereof, to determined if the combination antibodies, or antigen binding portions thereof have the predetermined target activity or a partial target activity;
- k) if steps h) or j) do not result in an antibody or antigen binding portion thereof having the predetermined target activity, or result in an antibody with only a partial activity, the method further comprising mutating additional amino acid residues selected from the group consisting of H33B, H52B and L31A to at least two other amino acid residues to thereby create a third panel of mutated antibodies or antigen binding portions thereof;
- l) evaluating the activity of the third panel of mutated antibodies or antigen binding portions thereof, to determine if a mutation of a single amino acid residue selected from the group consisting of H33B, H52B and L31A resulted in an antibody or antigen binding portion thereof, having the predetermined target activity or a partial activity;

- m) combining in a stepwise fashion in the parent antibody, or antigen binding portion thereof, individual mutation of step k) shown to have an improved activity, to form combination antibodies, or antigen binding portions, thereof;
- n) evaluating the activity of the combination antibodies or antigenbinding portions thereof, to determine if the combination antibodies, or antigen binding portions thereof have the predetermined target activity to thereby produce an antibody or antigen binding portion thereof with a predetermined target activity.
- A method for improving the activity of an antibody, or antigen-binding 101. portion thereof, comprising: 10
 - a) providing a parent antibody or antigen-binding portion thereof;

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- b) selecting a preferred selective mutagenesis position, contact or hypermutation position within a complementarity determining region (CDR) for mutation, thereby identifying a selected preferred selective mutagenesis position, contact or hypermutation position;
- c) individually mutating said preferred selective mutagenesis position, contact or hypermutation position to at least two other amino acid residues to thereby create a panel of mutated antibodies, or antigen-binding portions thereof;
- d) evaluating the activity of the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof;
- e) repeating steps b) through d) for at least one other preferred selective mutagenesis position, contact or hypermutation position if the desired antibody activity is not obtained;
- f) combining in a stepwise fashion, in the parent antibody, or antigen-binding portion thereof, individual mutations shown to have improved activity, to form combination antibodies, or antigen-binding portions thereof; and
 - g) evaluating the activity of the combination antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof; until an antibody, or antigen-binding portion thereof, with an improved activity, relative to the parent antibody, or antigen-binding portion thereof, is obtained.

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102. The method of claim 101, wherein contact positions are selected from the group consisting of H30, H31, H31B, H32, H33, H35, H50, H52, H52A, H53, H54, H56, H58, H95, H96, H97, H98, H101, L30, L31, L32, L34, L50, L52, L53, L55, L91, L92, L93, L94 and L96.

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- 103. The method of claim 101, wherein hypermutation positions are selected from the group consisting of H30, H31, H31B, H32, H52, H56, H58, L30, L31, L32, L53 and L93.
- 104. The method of claim 101, wherein the preferred positions are selected from the group consisting of H30, H31, H31B, H32, H33, H52, H56, H58, L30, L31, L32, L50, L91, L92, L93, L94.
- 105. The method of claim 101, wherein the contact positions are selected from the group consisting of L50 and L94.
 - 106. A method for improving the activity of an antibody, or antigen-binding portion thereof, comprising:
 - a) providing a recombinant parent antibody or antigen-binding portion thereof; that was obtained by selection in a phage-display system but whose activity is not further improved by mutagenesis in said phage-display system;
 - b) selecting a preferred selective mutagenesis position, contact or hypermutation position within a complementarity determining region (CDR) for mutation, thereby identifying a selected preferred selective mutagenesis position, contact or hypermutation position;
 - c) individually mutating said selected preferred selective mutagenesis position, contact or hypermutation position to at least two other amino acid residues to thereby create a panel of mutated antibodies, or antigen-binding portions thereof, and expressing said panel in a non-phage display system;
 - d) evaluating the activity of the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof;

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- e) repeating steps b) through d) for at least one other preferred selective mutagenesis position, contact or hypermutation position if the desired antibody activity is not obtained;
- f) combining, in the parent antibody, or antigen-binding portion thereof, individual mutations shown to have improved activity, to form combination antibodies, or antigen-binding portions thereof; and
 - g) evaluating the activity of the combination antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof; until an antibody, or antigen-binding portion thereof, with an improved activity, relative to the parent antibody, or antigen-binding portion thereof, is obtained.
 - 107. The method of claim 106, wherein contact positions are selected from the group consisting of H30, H31, H31B, H32, H33, H35, H50, H52, H52A, H53, H54, H56, H58, H95, H96, H97, H98, H101, L30, L31, L32, L34, L50, L52, L53, L55, L91, L92, L93, L94 and L96.
 - 108. The method of claim 106, wherein hypermutation positions are selected from the group consisting of H30, H31, H31B, H32, H52, H56, H58, L30, L31, L32, L53 and L93.

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- 109. The method of claim 106, wherein preferred selective mutagenesis positions are selected from the group consisting of H30, H31, H31B, H32, H33, H52, H56, H58, L30, L31, L32, L50, L91, L92, L93, L94
- The method of claim 106, wherein the contact positions are selected from the group consisting of L50 and L94.

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- 111. A method for improving the activity of an antibody, or antigen-binding portion thereof, comprising:
 - a) providing a recombinant parent antibody or antigen-binding portion thereof;
- b) selecting a preferred selective mutagenesis position, contact or hypermutation position within a complementarity determining region (CDR) for mutation, thereby identifying a selected preferred selective mutagenesis position, contact or hypermutation position;
 - c) individually mutating said selected preferred selective mutagenesis position, contact or hypermutation position to at least two other amino acid residues to thereby create a panel of mutated antibodies, or antigen-binding portions thereof and expressing said panel in an appropriate expression system;
 - d) evaluating the activity of the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof thereby identifying an activity enhancing amino acid residue;
- e) evaluating the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof, for at least one other property or characteristic, wherein the property or characteristic is one that needs to be retained in the antibody; until an antibody, or antigen-binding portion thereof, with an improved activity and at least one retained property or characteristic, relative to the parent antibody, or antigen-binding portion thereof, is obtained.
- 112. The method of claim 111, wherein contact positions are selected from the group consisting of H30, H31, H31B, H32, H33, H35, H50, H52, H52A, H53, H54, H56, H58, H95, H96, H97, H98, H101, L30, L31, L32, L34, L50, L52, L53, L55, L91, L92, L93, L94 and L96, and wherein the other property or characteristic is selected from the group consisting of preservation of non-crossreactivity with other proteins, preservation of non-crossreactivity with other human tissues, preservation of epitope recognition and an antibody with a close to germline immunoglobulin sequence.

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- selected from the group consisting of H30, H31, H31B, H32, H52, H56, H58, L30, L31, L32, L53 and L93, and wherein the other property or characteristic is selected from the group consisting of preservation of non-crossreactivity with other proteins, preservation of non-crossreactivity with other human tissues, preservation of epitope recognition and an antibody with a close to germline immunoglobulin sequence.
- 114. The method of claim 111, wherein the preferred selective mutagenesis positions are selected from the group consisting of H30, H31, H31B, H32, H33, H52, H56, H58, L30, L31, L32, L50, L91, L92, L93 and L94, and wherein the other property or characteristic is selected from the group consisting of preservation of non-crossreactivity with other proteins, preservation of non-crossreactivity with other human tissues, preservation of epitope recognition and an antibody with a close to germline immunoglobulin sequence.

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- 115. The method of claim 111, wherein the contact positions are selected from the group consisting of L50 and L94, and wherein the other property or characteristic is selected from the group consisting of preservation of non-crossreactivity with other proteins, preservation of non-crossreactivity with other human tissues, preservation of epitope recognition and an antibody with a close to germline immunoglobulin sequence.
- 116. A method for improving the activity of an antibody, or antigen-binding portion thereof, comprising:
- a) providing a recombinant parent antibody or antigen-binding portion thereof; that was obtained by selection in a phage-display system but whose activity cannot be further improved by mutagenesis in said phage-display system;
 - b) selecting a preferred selective mutagenesis position, contact or hypermutation position within a complementarity determining region (CDR) for mutation, thereby identifying a selected preferred selective mutagenesis position, contact or hypermutation position;

- c) individually mutating said selected preferred selective mutagenesis position, contact or hypermutation position to at least two other amino acid residues to thereby create a panel of mutated antibodies, or antigen-binding portions thereof, and expressing said panel in a non-phage display system;
- d) evaluating the activity of the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof thereby identifying an activity enhancing amino acid residue;
- e) evaluating the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof for at least one other property or characteristic, wherein the property or characteristic is one that needs to be retained, until an antibody, or antigen-binding portion thereof, with an improved activity and at least one retained property or characteristic, relative to the parent antibody, or antigen-binding portion thereof, is obtained.
- f) repeating steps a) through e) for at least one other preferred selective mutagenesis position, contact or hypermutation position;
- g) combining, in the parent antibody, or antigen-binding portion thereof, at least two individual activity enhancing amino acid residues shown to have improved activity and at least one retained property or characteristic, to form combination antibodies, or antigen-binding portions thereof; and
- h) evaluating the activity of the combination antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof; until an antibody, or antigen-binding portion thereof, with an improved activity and at least one retained property or characteristic, relative to the parent antibody, or antigen-binding portion thereof, is obtained.

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117. The method of claim 116, wherein contact positions are selected from the group consisting of H30, H31, H31B, H32, H33, H35, H50, H52, H52A, H53, H54, H56, H58, H95, H96, H97, H98, H101, L30, L31, L32, L34, L50, L52, L53, L55, L91, L92, L93, L94 and L96, and wherein the other property or characteristic is selected from the group consisting of preservation of non-crossreactivity with other proteins, preservation of non-crossreactivity with other human tissues, preservation of epitope recognition and an antibody with a close to germline immunoglobulin sequence.

- The method of claim 116, wherein the hypermutation positions are selected from the group consisting of H30, H31, H31B, H32, H52, H56, H58, L30, L31, L32, L53 and L93, and wherein the other property or characteristic is selected from the 5 group consisting of preservation of non-crossreactivity with other proteins, preservation of non-crossreactivity with other human tissues, preservation of epitope recognition and an antibody with a close to germline immunoglobulin sequence.
- The method of claim 116 wherein the preferred selective mutagenesis 119. positions are selected from the group consisting of H30, H31, H31B, H32, H33, H52, H56, H58, L30, L31, L32, L50, L91, L92, L93 and L94, and wherein the other property or characteristic is selected from the group consisting of preservation of noncrossreactivity with other proteins, preservation of non-crossreactivity with other human tissues, preservation of epitope recognition and an antibody with a close to germline 15 immunoglobulin sequence.
 - The method of claim 116, wherein the contact positions are selected from 120. the group consisting of L50 and L94, and wherein the other property or characteristic is selected from the group consisting of preservation of non-crossreactivity with other proteins, preservation of non-crossreactivity with other human tissues, preservation of epitope recognition and an antibody with a close to germline immunoglobulin sequence.
 - A method for improving the activity of an antibody, or antigen-binding portion thereof, comprising:
 - a) providing a recombinant parent antibody or antigen-binding portion thereof; that was obtained by selection in a phage-display system but whose activity cannot be further improved by mutagenesis in said phage-display system;

- b) selecting a contact or hypermutation position within a complementarity determining region (CDR) for mutation, thereby identifying a selected contact or hypermutation position;
- c) individually mutating said selected contact or hypermutation position to at least two other amino acid residues to thereby create a panel of mutated antibodies, or

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antigen-binding portions thereof, and expressing said panel in a non-phage display system;

- d) evaluating the activity of the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof thereby identifying an activity enhancing amino acid residue;
- e) evaluating the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof for at least one additional property or characteristic, wherein the property or characteristic is one that needs to be retained,
- until an antibody, or antigen-binding portion thereof, with an improved activity and at least one retained property or characteristic relative to the parent antibody, or antigen-binding portion thereof, is obtained.
- 122. The method of claim 121, wherein contact positions are selected from the group consisting of H30, H31, H31B, H32, H33, H35, H50, H52, H52A, H53, H54, H56, H58, H95, H96, H97, H98, H101, L30, L31, L32, L34, L50, L52, L53, L55, L91, L92, L93, L94 and L96, and wherein the other property or characteristic is selected from the group consisting of preservation of non-crossreactivity with other proteins, preservation of non-crossreactivity with other human tissues, preservation of epitope recognition and an antibody with a close to germline immunoglobulin sequence.
 - selected from the group consisting of H30, H31, H31B, H32, H52, H56, H58, L30, L31, L32, L53 and L93, and wherein the other property or characteristic is selected from the group consisting of preservation of non-crossreactivity with other proteins, preservation of non-crossreactivity with other human tissues, preservation of epitope recognition and an antibody with a close to germline immunoglobulin sequence.

- The method of claim 121, wherein the contact positions are selected from 124. the group consisting of L50 and L94, and wherein the other property or characteristic is selected from the group consisting of preservation of non-crossreactivity with other proteins, preservation of non-crossreactivity with other human tissues, preservation of epitope recognition and an antibody with a close to germline immunoglobulin sequence.
- A method for improving the activity of an antibody, or antigen-binding 125. portion thereof, comprising:
- a) providing a recombinant parent antibody or antigen-binding portion thereof; that was obtained by selection in a phage-display system but whose activity cannot be further improved by mutagenesis in said phage-display system;

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- b) selecting a preferred selective mutagenesis position, contact or hypermutation position within a complementarity determining region (CDR) for mutation, thereby identifying a selected preferred selective mutagenesis position contact or hypermutation position;
- c) individually mutating said selected preferred selective mutagenesis position. contact or hypermutation position to at least two other amino acid residues to thereby create a panel of mutated antibodies, or antigen-binding portions thereof, and expressing said panel in a non-phage display system;
- d) evaluating the activity of the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof thereby identifying an activity enhancing amino acid residue;
- e) evaluating the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof for at least one other property or characteristic, wherein the property or characteristic is one that needs to be retained, until an antibody, or antigen-binding portion thereof, with an improved activity and at least one retained property or characteristic relative to the parent antibody, or antigen-binding portion thereof, is obtained.
- f) repeating steps a) through e) for at least one other preferred selective mutagenesis position, contact or hypermutation position;
- g) combining, in the parent antibody, or antigen-binding portion thereof, at least two individual activity enhancing amino acid residues shown to have improved activity

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and at least one retained property or characteristic, to form combination antibodies, or antigen-binding portions thereof; and

- h) evaluating the activity of the combination antibodies. or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof; until an antibody, or antigen-binding portion thereof, with an improved activity and at least one retained property or characteristic relative to the parent antibody, or antigen-binding portion thereof, is obtained.
- 126. The method of claim 125, wherein contact positions are selected from the group consisting of H30, H31, H31B, H32, H33, H35, H50, H52, H52A, H53, H54, H56, H58, H95, H96, H97, H98, H101, L30, L31, L32, L34, L50, L52, L53, L55, L91, L92, L93, L94 and L96, and wherein the other property or characteristic is selected from the group consisting of preservation of non-crossreactivity with other proteins, preservation of non-crossreactivity with other human tissues, preservation of epitope recognition and an antibody with a close to germline immunoglobulin sequence.
 - selected from the group consisting of H30, H31, H31B, H32, H52, H56, H58, L30, L31, L32, L53 and L93, and wherein the other property or characteristic is selected from the group consisting of preservation of non-crossreactivity with other proteins, preservation of non-crossreactivity with other human tissues, preservation of epitope recognition and an antibody with a close to germline immunoglobulin sequence.

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positions are selected from the group consisting of H30, H31, H31B, H32, H33, H52, H56, H58, L30, L31, L32, L50, L91, L92, L93 and L94, and wherein the other property or characteristic is selected from the group consisting of preservation of non-crossreactivity with other proteins, preservation of non-crossreactivity with other human tissues, preservation of epitope recognition and an antibody with a close to germline immunoglobulin sequence.

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- 129. The method of claim 125, wherein the contact positions are selected from the group consisting of L50 and L94, and wherein the other property or characteristic is selected from the group consisting of preservation of non-crossreactivity with other proteins, preservation of non-crossreactivity with other human tissues, preservation of epitope recognition and an antibody with a close to germline immunoglobulin sequence.
- 130. A method for improving the activity of an antibody, or antigen-binding portion thereof, comprising:
 - a) providing a parent antibody or antigen-binding portion thereof;
- b) selecting a amino acid residue within a complementarity determining region (CDR) for mutation at a position other than H30, H31, H31B, H32, H33, H35, H50, H52, H52A, H53, H54, H56, H58, H95, H96, H97, H98, H101, L30, L31, L32, L34, L50, L52, L53, L55, L91, L92, L93, L94 and L96;
- c) individually mutating said selected position to at least two other amino acid
 residues to thereby create a panel of mutated antibodies, or antigen-binding portions thereof;
 - d) evaluating the activity of the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof thereby identifying an activity enhancing amino acid residue;
 - e) evaluating the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof, for changes in at least one other property or characteristic; until an antibody, or antigen-binding portion thereof, with an improved activity, relative to the parent antibody, or antigen-binding portion thereof, is obtained.

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- 131. A method for improving the activity of an antibody, or antigen-binding portion thereof, comprising:
 - a) providing a parent antibody or antigen-binding portion thereof;
- b) selecting a amino acid residue within a complementarity determining region (CDR) for mutation at a position other than H30, H31, H31B, H32, H33, H35, H50, H52, H52A, H53, H54, H56, H58, H95, H96, H97, H98, H101, L30, L31, L32, L34, L50, L52, L53, L55, L91, L92, L93, L94 and L96;

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c) individually mutating said selected position to at least two other amino acid residues to thereby create a panel of mutated antibodies, or antigen-binding portions thereof;

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- d) evaluating the activity of the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof, thereby identifying an activity enhancing amino acid residue;
 - e) repeating steps b) through d) for at least one other position within the CDR which is neither the position selected under b) nor a position at H30, H31, H31B, H32, H33, H35, H50, H52, H52A, H53, H54, H56, H58, H95, H96, H97, H98, H101, L30, L31, L32, L34, L50, L52, L53, L55, L91, L92, L93, L94 and L96;

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- f) combining, in the parent antibody, or antigen-binding portion thereof, at least two individual activity enhancing amino acid residues shown to have improved activity, to form combination antibodies, or antigen-binding portions thereof; and
- g) evaluating the activity of the combination antibodies, or antigen-binding portions thereof with two activity enhancing amino acid residues, relative to the parent antibody or antigen-binding portion thereof until an antibody, or antigen-binding portion thereof, with an improved activity, relative to the parent antibody, or antigen-binding portion thereof, is obtained.
- A method for improving the activity of an antibody, or antigen-binding 20 132. portion thereof, comprising:
 - a) providing a recombinant parent antibody or antigen-binding portion thereof; that was obtained by selection in a phage-display system but whose activity cannot be further improved by mutagenesis in said phage-display system;
 - b) selecting a selecting an amino acid residue within a complementarity determining region (CDR) for mutation at a position other than H30, H31, H31B, H32, H33, H35, H50, H52, H52A, H53, H54, H56, H58, H95, H96, H97, H98, H101, L30, L31, L32, L34, L50, L52, L53, L55, L91, L92, L93, L94 and;
- c) individually mutating said selected contact or hypermutation position to at least two other amino acid residues to thereby create a panel of mutated antibodies, or 30 antigen-binding portions thereof, and expressing said panel in a non-phage display system;

- d) evaluating the activity of the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof thereby identifying an activity enhancing amino acid residue;
- e) evaluating the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof, for changes in at least one other property or characteristic, until an antibody, or antigen-binding portion thereof, with an improved activity, relative to the parent antibody, or antigen-binding portion thereof, is obtained.
- A method for improving the activity of an antibody, or antigen-binding 133. portion thereof, comprising:

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- a) providing a parent antibody or antigen-binding portion thereof that was obtained by selection in a phage-display system but whose activity cannot be further improved by mutagenesis in said phage-display system;
- b) selecting an amino acid residue within a complementarity determining region (CDR) for mutation at a position other than H30, H31, H31B, H32, H33, H35, H50, H52, H52A, H53, H54, H56, H58, H95, H96, H97, H98, H101, L30, L31, L32, L34, L50, L52, L53, L55, L91, L92, L93, L94 and L96;
- c) individually mutating said selected position to at least two other amino acid residues to thereby create a panel of mutated antibodies, or antigen-binding portions thereof and expression in a non-phage display system;
- d) evaluating the activity of the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof thereby identifying an activity enhancing amino acid residue;
- e) repeating steps b) through d) for at least one other position within the CDR which is neither the position selected under b) nor a position at H30, H31, H31B, H32, H33, H35, H50, H52, H52A, H53, H54, H56, H58, H95, H96, H97, H98, H101, L30, L31, L32, L34, L50, L52, L53, L55, L91, L92, L93, L94;
- f) combining, in the parent antibody, or antigen-binding portion thereof, at least two individual activity enhancing amino acid residues shown to have improved activity, to form combination antibodies, or antigen-binding portions thereof; and

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g) evaluating the activity and other property or characteristics of the combination antibodies, or antigen-binding portions thereof, with two activity enhancing amino acid residues, relative to the parent antibody or antigen-binding portion thereof; until an antibody, or antigen-binding portion thereof, with an improved activity, relative to the parent antibody, or antigen-binding portion thereof, is obtained.

- 134. A method for improving the activity of an antibody, or antigen-binding portion thereof, without affecting other properties, comprising:
 - a) providing a parent antibody or antigen-binding portion thereof;

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- b) selecting an amino acid residue within a complementarity determining region (CDR) for mutation at a position other than H30, H31, H31B, H32, H33, H35, H50, H52, H52A, H53, H54, H56, H58, H95, H96, H97, H98, H101, L30, L31, L32, L34, L50, L52, L53, L55, L91, L92, L93, L94 and L96;
- c) individually mutating said selected position to at least two other amino acid residues to thereby create a panel of mutated antibodies, or antigen-binding portions thereof:
 - d) evaluating the activity of the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof thereby identifying an activity enhancing amino acid residue;
 - e) evaluating the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof, for changes in at least one other property or characteristic, wherein the property or characteristic needs to be retained, until an antibody, or antigen-binding portion thereof, with an improved activity and retained property, or characteristic relative to the parent antibody, or antigen-binding portion thereof, is obtained.
 - 135. A method for improving the activity of an antibody, or antigen-binding portion thereof, comprising:
 - a) providing a parent antibody or antigen-binding portion thereof;
 - b) selecting an amino acid residue within a complementarity determining region (CDR) for mutation at a position other than H30, H31, H31B, H32, H33, H35, H50,

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- H52, H52A, H53, H54, H56, H58, H95, H96, H97, H98, H101, L30, L31, L32, L34, L50, L52, L53, L55, L91, L92, L93, L94 and L96;
- c) individually mutating said selected position to at least two other amino acid residues to thereby create a panel of mutated antibodies, or antigen-binding portions thereof;
- d) evaluating the activity of the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof, thereby identifying an activity enhancing amino acid residue;
- e) evaluating the panel of mutated antibodies or antigen-binding portions thereof, relative to the parent antibody or antigen-portion thereof, for changes in at least one other property or characteristic;
 - f) repeating steps b) through e) for at least one other CDR position which is neither the position selected under b) nor a position at H30, H31, H31B, H32, H33, H35, H50, H52, H52A, H53, H54, H56, H58, H95, H96, H97, H98. H101, L30, L31, L32, L34, L50, L52, L53, L55, L91, L92, L93, L94 and L96;
 - g) combining, in the parent antibody, or antigen-binding portion thereof, at least two individual activity enhancing amino acid residues shown to have improved activity but not affecting at least one other property or characteristic, to form combination antibodies, or antigen-binding portions thereof with at least one retained property or characteristic; and
 - h) evaluating the activity and the retention of at least one property of characteristic of the combination antibodies, or antigen-binding portions thereof with two activity enhancing amino acid residues, relative to the parent antibody or antigen-binding portion thereof until an antibody, or antigen-binding portion thereof, with an improved activity and at least one retained property or characteristic, relative to the parent antibody, or antigen-binding portion thereof, is obtained.
 - 136. A method to improve the affinity of an antibody or antigen-binding portion thereof, comprising:
- a) providing a parent antibody or antigen-binding portion thereof that was obtained by selection in a phage-display system but whose activity cannot be further improved by mutagenesis in said phage-display system;

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- b) selecting an amino acid residue within a complementarity determining region (CDR) for mutation other than H30, H31, H31B, H32, H33, H35, H50, H52, H52A, H53, H54, H56, H58, H95, H96, H97, H98, H101, L30, L31, L32, L34, L50, L52, L53, L55, L91, L92, L93, L94 and L96;
- c) individually mutating said selected position to at least two other amino acid residues to thereby create a panel of mutated antibodies, or antigen-binding portions thereof and expression in a non-phage display system;
- d) evaluating the activity of the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof thereby identifying an activity enhancing amino acid residue;
- e) evaluating the panel of mutated antibodies, or antigen-binding portions thereof, relative to the parent antibody or antigen-binding portion thereof, for changes in at least one other property or characteristic until an antibody, or antigen-binding portion thereof, with an improved activity, relative to the parent antibody, or antigen-binding portion thereof, is obtained.
- 137. A method for improving the activity of an antibody, or antigen-binding portion thereof, comprising:
- a) providing a parent antibody or antigen-binding portion thereof that was obtained by selection in a phage-display system but whose activity cannot be further improved by mutagenesis in said phage-display system;
- b) selecting an amino acid residue within a complementarity determining region (CDR) for mutation other than H30, H31, H31B, H32, H33, H35, H50, H52, H52A, H53, H54, H56, H58, H95, H96, H97, H98, H101, L30, L31, L32, L34, L50, L52, L53, L55, L91, L92, L93, L94 and L96;
- c) individually mutating said selected position to at least two other amino acid residues to thereby create a panel of mutated antibodies, or antigen-binding portions thereof and expression in a non-phage display system;
- d) evaluating the activity and retention of at least one other property or
 characteristic of the panel of mutated antibodies, or antigen-binding portions thereof,
 relative to the parent antibody or antigen-binding portion thereof, thereby identifying an activity enhancing amino acid residue;

- e) repeating steps b) through d) for at least one other CDR position which is neither the position selected under b) nor a position at H30, H31, H31B, H32, H33, H35, H50, H52, H52A, H53, H54, H56, H58, H95, H96, H97, H98, H101, L30, L31, L32, L34, L50, L52, L53, L55, L91, L92, L93, L94 and L96;
- f) combining, in the parent antibody, or antigen-binding portion thereof, at least two individual activity enhancing amino acid residues shown to have improved activity and not to affect at least one other property or characteristic, to form combination antibodies, or antigen-binding portions thereof; and
- g) evaluating the activity and retention of at least one other property or characteristic of the combination antibodies, or antigen-binding portions thereof with two activity enhancing amino acid residues, relative to the parent antibody or antigen-binding portion thereof until an antibody, or antigen-binding portion thereof, with an improved activity and at least one other retained property or characteristic, relative to the parent antibody, or antigen-binding portion thereof, is obtained.

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- 138. The method of claims 130, 131, 132, 133, 134, 135, 136 or 137, wherein the other property or characteristic is selected from the group consisting of preservation of non-crossreactivity with other proteins, preservation of non-cross reactivity with other human tissues. preservation of epitope recognition and an antibody with a close to germline immunoglobulin sequence.
- 139. A method for detecting human IL-12 comprising contacting human IL-12 with the antibody, or antigen-binding portion thereof, of any of claims 1-52, 74-83 and 87 such that human IL-12 is detected.

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- 140. The method of claim 139, wherein human IL-12 is detected in vitro.
- 141. The method of claim 139, wherein human IL-12 is detected in a biological sample for diagnostic purposes.

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142. Use of the antibody, or antigen-binding portion thereof, of any of claims 1-52, 74-83 and 87 in therapy.

143. Use of the antibody, or antigen-binding portion thereof, of any of claims 1-52, 74-83 and 87 in the manufacture of a medicament for the treatment of a a disorder in which IL-12 activity is detrimental.

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- 144. The use of claim 143, wherein the disorder is selected from the group consisting of rheumatoid arthritis, osteoarthritis, juvenile chronic arthritis, Lyme arthritis, psoriatic arthritis, reactive arthritis, spondyoarthropathy, ankylosing spondylitis, systemic lupus erythematosis, Crohn's disease, ulcerative colitis, inflammatory bowel disease, multiple sclerosis, insulin dependent diabetes mellitus, thyroiditis, asthma, allergic diseases, psoriasis, dermatitisscleroderma, thyroiditis, graft versus host disease, organ transplant rejection, acute or chronic immune disease associated with organ transplantation, sarcoidosis, atherosclerosis, disseminated intravascular coagulation, Kawasaki's disease, Grave's disease, nephrotic syndrome, chronic fatigue syndrome, polyarteritis nodosa, Wegener's granulomatosis, Henoch-Schonlein purpura, microscopic vasculitis of the kidneys, chronic active hepatitis, Sjogren's syndrome, uveitis, sepsis, septic shock, sepsis syndrome, adult respiratory distress syndrome, cachexia, infectious diseases, parasitic diseases, acquired immunodeficiency syndrome, acute transverse myelitis, myasthenia gravis, Huntington's chorea, Parkinson's disease, Alzheimer's disease, stroke, primary biliary cirrhosis. fibrotic lung diseases, hemolytic anemia, malignancies, heart failure and myocardial infarction.
 - 145. The use of claim 143, wherein the disorder is Crohn's disease.

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- 146. The use of claim 143, wherein the disorder is multiple sclerosis.
- 147. The use of claim 143, wherein the disorder is rheumatoid arthritis.

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56 55 54 53 52A 52 51 50 FIRYDGSN CDR H2 49 48 47 46 45 44 43 42 41 40 39 37 36 WVRQAPGKGLEWVA 35 34 33 32 31 SYGMH H CDR QVQLVQSGGGVVQPGRSLRLSCAASGFTF AO3 VH germline Y61 VH germine 136-15 VH germline Kabat number Y61 L50Y VH Y61-H31E VH 103-14 VH Cos-3/JH3 136-15 VH 101-11 VH 149-6 VH 103-4 VH 103-8 VH 149-5 VH 78-34 VH Y139 VH 26-1 VH 70-1 VH 79-1 VH Y61 VH AO3 VH G6 VH 69 23 39 43 45 49 53 55 57 59 19 63 65 6 7 33 5 41 51 47

Figure 1A. Heavy Chain Variable Region Sequences

Figure 1B. Heavy Chain Variable Region Sequences

CDR H3	113 112 111 110 109 108 107 106 105 104 103 102 101 98 97 95	SGSYDY WGQGTMVTVSS					нн.х	нн.х	нн.х	N.HH	HH.T		H H	I	. H. H. N	и.н	N.HH .	N.HH .	N.HH .	•	H	H	N . H		N.HH .
	943 921 998 887 882 882 882 8777777777777777777	RFTISRDNSKNTLYLOMKSLRAEDTAVYYCTT	A						× · · · · · · · · · · · · · · · · · · ·	X				× · · · · · · · · · · · · · · · · · · ·	Y	× · · · · · · · · · · · · · · · · · · ·	× · · · · · · · · · · · · · · · · · · ·	× · · · · · · · · · · · · · · · · · · ·	ж	M · · · · · · · · · · · · · · · · · · ·	ж	M · · · · · · · · · · · · · · · · · · ·	× · · · · · · · · · · · · · · · · · · ·	× · · · · · · · · · · · · · · · · · · ·	X
CDR H2	Kabat number 28699999	JOE9wt VH KYYADSVKG	Cos-3/JH3 VH	70-1 VH	78-34 VH	79-1 VН	101-11 VH	26-1 VH	136-15 VH	136-15 VH germline	149-5 VH	149-6 VH	103-4 VH	103-8 VH	103-14 VH	G6 VH	Y139 VH	AO3 VH	AO3 VH germline	у61 vн	Y61 VH germine	Y61-H31E VH	Y61 L50Y VH	Y61-194Y VH	7695
	SEQ ID NO:	33 JC	35 Cc	37 76	39 76	41 79	43 10	45 20	47 1.	49 13	51 1	53 1	55 1	57 1	59 1	61 G	63 Y	65 A	67 A	23 Y	¥ 69	71 Y	73 Y	75 Y	31 5

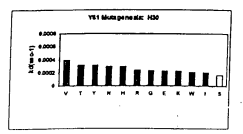
	CDR L2	5 6 5 5 5 4 5 3 5 2 5 1 5 0 4 9 4 8	Y GNDQRP	S N															•				· · · · · · · · · · · · · · · · · · ·		· · · · · · · · · · · · · · · · · · ·
ses.		47 46 45 44 43 42 41 40 39 38 37 36	YQQLPGTAPKLL				•												•						•
Variable Region Sequences	CDR L1	34 33 32 31 30 28 27 27 27 27 27 26 25	GGRSNIGSNTV	T.SSAGYD.H					· · · · · · · · · · · · · · · · · · ·										· · · · · · · · · · · · · · · · · · ·					· · · · · · · · · · · · · · · · · · ·	
Figure 1C. Light Chain Vari		23 22 21 20 19 115 115 114 132 111 98 87 76 55 44 32 21	PSVSGTPGQRVTIS	OSA															OS		SO		S		OS
		Kabat number	Joe9 VL wt	DP18 Lv1042/JA1	78-34 VL	79-1 VL	101-11 VL	26-1 VL	S	136-15 VL germline	149-5 VL	149-6 VL	103-4 VL	103-8 VL	103-14 VL	TA 95	X139 VL	AO3 VL	AO3 VL germline	Y61 VL	Y61 VL germline	Y61-H31E VL	Y61-L50Y VL	X61-L94Y VL	J695 VL
		SEQ ID NO:	34	38	40	42	44	46	48	20	52	54	26	28	9	62	64	99	99	24	70	72	74	16	32

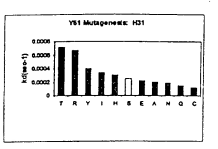
Figure 1D. Light Chain Variable Region Sequences

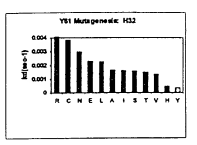
		GTKVTVL			•	•	•	•		•	•			•										• • • • • • • • • • • • • • • • • • • •	
CDR L3	950 958 958 958 94 93 93	DSSLRGSR	,		RGFT	· · · · · · · · · · · · · · · · · · ·	RGFT	٠	KGF	KGF	•	GFT.	GFT.A	G F	. ERGFT M	GTHPLT	GSHP	GTHPLT	GTHPLT	GTHPAL	GTHPAL	RGTHPALL	GTHPAL	YTHPAL	
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	Kabat number	Joe9 VL wt	Dp18 Lv1042/JA1	70-1 VL	78-34 VL	79-1 VL	101-11 VL	26-1 VL	136-15 VL	136-15 VL germline	149-5 VL	149-6 VL	103-4 VL	103-8 VL	103-14 VL	G6 VL	Y139 VL	AO3 VL	AO3 VL germline	Y61 VL	Y61 VL germline	Y61-H31E VL	X61-L50Y VL	Y61-L94Y VL	J695 VL
_	SEQ ID NO:	34	36	38	40	42	44	46	48	20	52	54	26	28	09	62	64	99	89	24	70	72	74	76	32

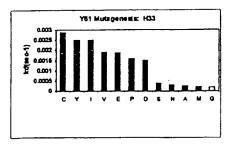
Figure 2A. Y61 Heavy Chain CDR H1 Mutagenesis

SEQ ID NO: 2 8 9 0 1 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3 3	34	يبا	korr
			(x 10 ⁵)
288 E	М	Н	L
			22.8
289 S		1.	16.8
290 Y			31.9
291 H	•		29.6
292 K			22.5
293 R			24.5
294 N			30.1
295 T	.]		32.0
296 G	$\overline{\cdot}$		23.3
297 V	.		39.9
298 I	$\overline{\cdot}$		20.7
299 W	$\overline{\cdot}$		21.6
300 E	$\overline{\cdot}$		21.9
301 C		. 1	12.0
302	-	-1	24.9
303 Y		╗	39.8
304 H	. 1	$\overline{\cdot}$	30.9
305 R		\exists	66.4
306 N	$\overline{\cdot}$	$\overline{\cdot}$	19.1
307 Q		-	15.2
308 T	.	ᄀ	71.6
309 A	.	. 1	20.5
310		ᄀ	33.4
311 E .	. 1	$\overline{\cdot}$	229.0
312			383.0
313	- 1	•	157.5
314 Y .	. 1		33.7
315 H .	. 1		46.1
316 R .	.		448.5
317 N .			297.0
318 T .			148.0
319 A .	. 1		165.5
320 V .	. 1		133.5
321 L .	\neg	$\overline{\cdot}$	226.0
322 I .		$\overline{\cdot}$	160.5
323 D	$\overline{\cdot}$	$\overline{\cdot}$	152.0
324 E	.		189.0
325 C			286.5
326		\Box	39.9
327 Y	. 1	.	250.5
328 N	. 1	\exists	30.8
329 G	. 1	. 1	17.8
330 A			27.3
331 v	.		191.0
332 M		. 1	21.5
333 I	•		250.0
334 P	. 1		159.5





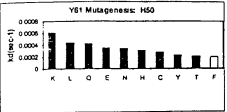


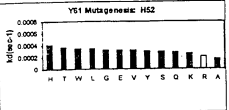


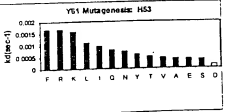
6/14

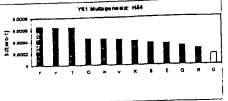
Figure 2B. Y61 Heavy Chain CDR H2 Mutagenesis

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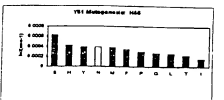




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Figure 2C. Y61 Heavy Chain CDR H2 Mutagenesis

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		L	CDR H2																
SEQ ID NO:		50	51	52	52A	53	54		56	57	85	59	69	61	62	63	62	65	k _{err} (x 10 ⁵)
19	Y61	F	Ī	R	Y	D	G	s	N	K	Y	Y	Α	D	S	V	К	G	
383		\cdot		Ŀ		Ŀ	£	Ŀ	Ŀ	Ŀ	Ŀ	·	$\overline{}$			\cdot	<u>.</u>	·	66.3
384			•		·	٠	·	•	S		Ŀ	·	•		ك	·	·	\cdot	62.4
385			\cdot	$ \cdot $		٠,		٠	Y		$ \cdot $	\cdot		٠i	<u>. </u>	╛	٠	\cdot	39.0
386		•	٠	·			•	•	Н		٠	\cdot	\cdot	•]	•		\cdot	اــــــــــــــــــــــــــــــــــــــ	42.0
387			•		•	•			N	•			•	.]	·	·	.	$\cdot \mid$	38.5
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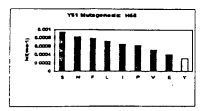
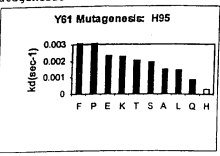
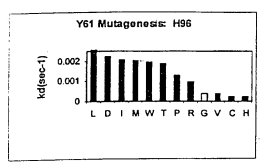
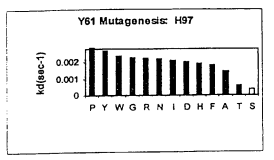


Figure 2D. Y61 Heavy Chain CDR H3 Mutagenesis

		1						
				Т	Т	Τ	Τ	,
SEQ ID		95	96	97	98	101	102	(x 10 ⁵)
17	Y61	Н	G	S	Н	D	N	
404		E		1	١.		1.	231.5
405		s	·		T .	١.	T.	193.0
406		Н	·		1.			28.7
407		К	١.	·	Ι.	1.		227.5
40B		Q	$\overline{}$	Γ.				85.9
409		T						202.0
410		Α						150.0
411		L						147.5
412	-	P	•			$\overline{}$		471.0
413		F	•				. 1	514.0
414			D					223.5
415			c		•	•		24.2
416		•	Н	÷.				23.7
417			R					96.2
418		Ť	T					186.0
419			G	·	÷			39.7
420			v	Ť	÷			38.2
421			М			÷		204.5
422		÷	L	÷	\div	\exists	·	261.0
423		$\dot{\cdot}$	Ĩ		Ħ			207.5
424	_		P					129.0
425			W	•	·	-	-	197.0
426		÷	. 	$\overline{}$	·			202.0
427					•			37.5
428				_	\neg			273.0
429		•		H		\vdots		190.5
430				R	÷			224.0
431		Ħ		N				221.5
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434		Ť		A				143.0
435		-		Ī	·	÷		208.0
436			•	P	$\overline{\cdot}$			300.0
437				W				239.0
438		$\overline{}$		F				180.5
439				-	Н			25.5
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447		÷			-	D		14.4
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449					·	Y	÷	465.0
450					÷	H		327.0
451					•	R		110.0







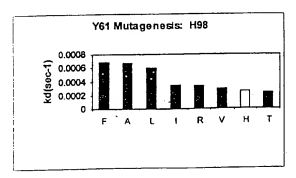
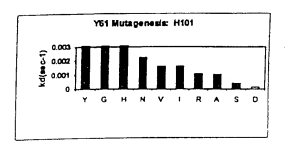


Figure 2E. Y61 Heavy Chain CDR H3 Mutagenesis

	Γ	Г						
SEQ ID		95	96	97	98	101	102	K _{off} (x 10 ⁵)
17	Y61	Н	G	S	Н	D	N	
452						7		223.0
453						G		375.0
454					\cdot	Α		106.5
455						٧	·	163.0
456					•	I		162.5
457		$\overline{\cdot}$				•	S	32.5
458		$\overline{\cdot}$				•	Н	18.0
459		$\overline{\cdot}$		•	.]	•	К	40.5
460					\cdot	•	R	57.5
461		- 1		•		•	2	40.3
462				.			T	33.3
463			•				G	69.2
464			•	. 1			A	38.2
465		$\overline{\cdot}$					L	95.6
466			$\overline{\cdot}$. 1		•	Ι	99.6
467						•	Р	181.5
468			-			•	W	23.5
469			$\overline{\cdot}$				F	31.8



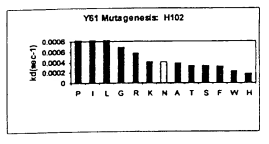
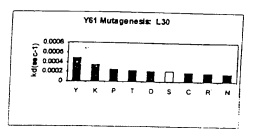
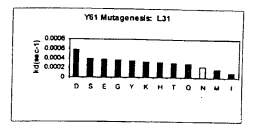


Figure 2F. Y61 Light Chain CDR L1 Mutagenesis

		_								_	_	_						
	-	╁	_	т-		_		20	<u> </u>	L)	-	_	_			_		_
SEQ ID NO:		24	25	26	12		3 7	27B	28	29	٤	3	ا ا	32	3	34	k _{ott} (x 10	5)
22	Y61	s	G	G	R	S	1	V	I	G	s	1	1	Т	v	к		_
470		ŀ	Ŀ	Ŀ	Ŀ	Ŀ		. [•	D						22.0)
471		<u> </u>			•		Γ.	. [•	С	Τ.	Т	.]		٠	18.6	5
472		Ŀ				Ŀ	Τ.		.]		s	1.	T	. 1	$\overline{\cdot}$		21.1	_
473		٠			•		Τ.	7	. 1	$\overline{\cdot}$	Y	١.	1	. 1	. 1	$\overline{\cdot}$	48.3	;
474							1.	T	. 1	.	K	٦.	T	.	.	.	34.6	
475		•		\cdot		$\overline{}$	1.	T	. T		R		T	. 1	.	.	18.2	
476							١.	Τ,	1	.	N	١.	Τ	. T	.	. 1	16.6	
477	_	$\overline{}$]		.]		1.	Τ.		.	T		1.	. T		.	22.6	
478		\cdot	$\overline{\cdot}$	\cdot				٦.	T	$\overline{\cdot}$	P	Ī.	Τ.	1	.	.	25.0	
479		\cdot	$\overline{\cdot}$					Τ.	T	. 1		D	١.		.	.	58.0	
480		\cdot	\cdot	\cdot	\cdot		Γ.	٦.	T	. 1		E	١.	1	.	\Box	38.4	٦
481		\cdot	\cdot					٦.	Т	. T		s	١.	Т	.	.	39.2	
482		$\cdot $. [Τ.	Τ	. 1		Y	١.	T	.	. T	35.7	٦
483		\cdot	\cdot	T	.]			٦.	T	. T		Н	١.	Τ.	. T	.	31.5	٦
484		$\overline{\cdot}$.				١.	1	.		к	١.	Τ.	. 1	.	33.1	٦
485		\cdot	\cdot	.				١.	T	. T	. 1	N	Γ.	Τ.	.	.	22.9	٦
486		\cdot		. [.			1.	T	.	. 1	Q		Τ.	.	.	29.2	٦
487			. [.	. 1				Τ.	.	. 1	T		٦.		.	30.9	٦
488		\cdot	\cdot	. [.]		•	١.	Τ.	T		G		Τ.	Π.	. 1	36.6	٦
489		\cdot	. [.					1	.	. 1	M		١.	Τ.	.	17.4	٦
490	T	\cdot	$\cdot \top$.	. [.	-	Ī .	T.	T	.	Ī	•	Τ.	١,	.	9.7	٦
491		$\cdot \mathbb{I}$	\cdot	\cdot	$\cdot \top$				Τ.	T	. 1		D	١.	Τ.	7	25.2	7
492		. [\cdot	. [. T			·	Τ.		.]	,	С	٦.	٦.	.	381.5	٦
493			. [$\cdot $	$\cdot \top$				Τ.	Т	.]		S	٦.	Τ.	.	191.0	7
494		$\cdot \mathbb{I}$	$\cdot $	$\cdot T$. [.]			Τ.	T	. [Y	١.	Τ.		21.3	7
495		٠I	\cdot	\cdot	\Box				٦.	Т			Н	Ι.	Τ.		26.0	
496	-1	. [. [Τ.	1	.]		K	٦.	7.	\Box	31.8	7
497			·I	. [\cdot			•	Τ.				R	1.	٦.		690.0]
498		. [\cdot	$\cdot \top$.	\Box			٦.	T	.]		N	١.	٦.	П	196.5	7
499		$\cdot \top$	$\cdot \top$	$\cdot \top$	$\cdot \top$.]			١.	T	. T		Q	Τ.	٦.	7	247.0	٦
500		. [.	\cdot	. 1	. 1	•		١.	_	. 1		T	Τ.	1.	1	24.1	٦
501					١.	Ť.	.		Ā	١.	٦.	7	190.5	٦
502	 1			1.	1	.		v	١.	٦.	\top	164.5	٦
503			.	_	_				١.	1	.		L	1.	1.	-	215.5	7
504		.	_	.	_	.		•	١.	Τ.	1		Ī	۲.	١.	1	154.0	7
505		\Box	.	.	. \dagger	.†	\exists		T.	+	.†	.	P	١.	Τ.	7	42.4	1





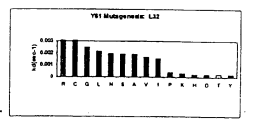
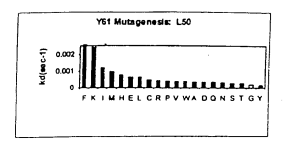


Figure 2G. Y61 Light Chain CDR L2 Mutagenesis

		L		CI	R				
SEQ ID NO:		50	51	52	53	54	55	56	
20	Y61	G	И	D	Q	R	P	S	_
506		D		•	•				34.8
507		Ξ				•	•		61.7
508		С						٠	46.7
509		S	\cdot	٠	•				28.6
510	•	Y	$\overline{\cdot}$		•		•		17.4
511		H	$\overline{\cdot}$						76.1
512		К		.	.				242.5
513		R							44.4
514		N				$\overline{\cdot}$.]	$\overline{\cdot}$	30.5
515		Q	.						34.8
516		T	.]	.	.]			\cdot	27.2
517		G							21.5
518		A	$\overline{\cdot}$	$\overline{\cdot}$	$\overline{\cdot}$	$\overline{\cdot}$			37.2
519		V	$\overline{\cdot}$.]	.			.	38.5
520		М	$\overline{\cdot}$	$\overline{\cdot}$	$\overline{\cdot}$				95.3
521		L					. [$\overline{\cdot}$	61.6
522	$\neg \neg$	I	\neg	.	$\overline{\cdot}$				120.5
523		P		$\overline{\cdot}$				$\overline{\cdot}$	41.0
524		W	. 1	.	ᄀ		\Box		38.2
525		F		$\overline{\cdot}$. 1		-1	$\overline{\cdot}$	3,476.7
526		. 1	.	. 1	s				86.6
527	1	$\overline{\cdot}$			Y			$\overline{\cdot}$	73.3
528					R		$\overline{\cdot}$		61.4
529	$\neg \neg$.]	Q	- 1	\Box	$\overline{\cdot}$	29.7
530			-		T				83.4
531	1		\cdot		Α	.			55.4
532			$\overline{\cdot}$		I			$\overline{\cdot}$	85.5
533		·	$\overline{\cdot}$		Р				97.4



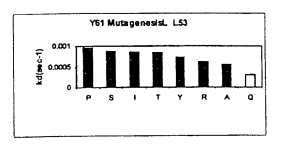
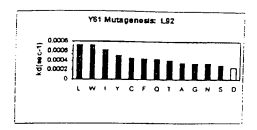
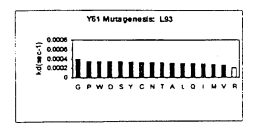


Figure 2H. Y61 Light Chain CDR L3 Mutagenesis

	1	CDR L3												
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18	Y61	Q	s	Y	D	R	G	T	Н	P	Α	L	L	
534		١.	1.	1.	D					Ŀ				25.9
535					С	Ī.		١.		1.				45.3
536					S		i .	ī.						30.7
537					Y			$\overline{}$	Ī.			, [51.1
538	-				N									34.7
539					Q					•	\cdot	\Box	$\overline{}$	42.7
540					T					$\overline{\cdot}$		\cdot	$\cdot \bot$	40.8
541					G							\cdot	. [34.9
542				$\overline{\cdot}$	Α	.	.	.]				$\overline{\cdot}$		35.7
543					L	$\overline{\cdot}$.]		\cdot		$\overline{\cdot}$.	72.8
544		\cdot		\cdot	I		. [$\overline{\cdot}$	\cdot		.]	\cdot	.	61.8
545				.]	W	$\overline{\cdot}$	$\overline{\cdot}$			•		.]	<u>.</u> ا	72.0
546		$\overline{\cdot}$	$\overline{\cdot}$		F	\cdot				$\overline{\cdot}$	$\overline{\cdot}$.	44.9
547			$\overline{\cdot}$			D		\Box		$\overline{\cdot}$	$\overline{\cdot}$. [$\overline{\cdot}$	34.3
548		$\overline{\cdot}$	$\overline{\cdot}$.	С	$\overline{\cdot}$.]		$\overline{\cdot}$. [32.0
549		. 1	$\overline{\cdot}$	$\overline{\cdot}$. 1	S	$\overline{\cdot}$. 1		.]	$\overline{\cdot}$	- [34.1
550				$\overline{\cdot}$	$\overline{\cdot}$	Y	<u>.</u> i	. 1	\Box	- 1			. [33.5
551		-1		$\overline{\cdot}$. 1	R	$\overline{\cdot}$.		.]	. [•	$\overline{\cdot}$	19.9
552		. 1	$\overline{\cdot}$. 1		N	.	\cdot	.		$\overline{\cdot}$	\cdot	\Box	31.6
553			$\overline{\cdot}$	$\overline{\cdot}$	$\overline{\cdot}$	Q		. 1	. 1	\Box	$\overline{\cdot}$	\cdot	$\overline{\cdot}$	30.0
554				.	.	T		. 1		. 1	$\overline{\cdot}$	$\cdot $	$\overline{\cdot}$	31.6
555			\Box	. 1		G		. 1	. 1		$\overline{\cdot}$	\cdot		39.2
556		. 1	\exists	$\overline{\cdot}$		A		\Box		$\overline{\cdot}$	\cdot		.]	31.0
557					. 1	V	. 1			$\overline{\cdot}$	\neg		$\overline{\cdot}$	26.9
558			\Box	.	. 1	м	. 1	$\overline{\cdot}$. 1	\Box			27.5
559			$\overline{\cdot}$. 1	L	$\overline{\cdot}$. 1		$\overline{\cdot}$	\Box			30.0
560		$\overline{\cdot}$	$\overline{\cdot}$. 1	I		\neg	•	$\overline{\cdot}$. 1	\exists	$\overline{\cdot}$	29.5
561		$\overline{\cdot}$. 1	P	. 1	$\overline{\cdot}$.	$\overline{\cdot}$.]	\cdot	34.9
562	-		•			W				.	$\overline{\cdot}$		\Box	34.9
563				. 1			Б	$\overline{\cdot}$			\exists	\cdot	\Box	25.3
564							c	-				_	.]	52.0
565					Ħ		s				$\overline{\cdot}$	$\overline{\cdot}$	\Box	28.7
566					$\overline{}$		Y				$\overline{}$		$\overline{\cdot}$	13.1
567						- 1	H					$\overline{}$	$\overline{\cdot}$	18.7
568					Ť	Ť	R	÷	•					23.1
569		Ť	÷		Ť		N	$\overline{}$	·			. 1	$\overline{}$	13.7
570		÷	÷	÷	Ť	Ì	0			•		. 1	. 1	25.0
571			<u>:</u>				Ť	÷		Ħ	Ť	. 1		30.5
572		·	÷	÷			Ġ	-		·		-1		25.6
573		÷			÷		A		÷	H.	-	÷	-	52.6
574		H		-	ᅴ		v			Ė		-1		35.1
575		H	\vdash	_	÷		L	·	÷	i.		ij		24.4
576		<u> </u>	\vdash		$\dot{-}$		ij	·	·	i -	÷	÷		27.6
577		·	-			·	P	÷	·	÷		$\dot{\exists}$	-	33.2
578		·	•	·	<u>-</u>	·	W	-	·	•		\vdots	-	29.3
579		\vdash	\vdash	\cdot		-	_	-	÷	\vdash	H	-	-	23.6
5/9		ك	۰	لنا	ا ن	لنا	F	ك	<u> </u>	٠.	ك	ك	. 1	





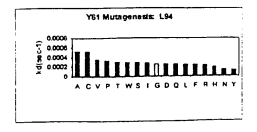
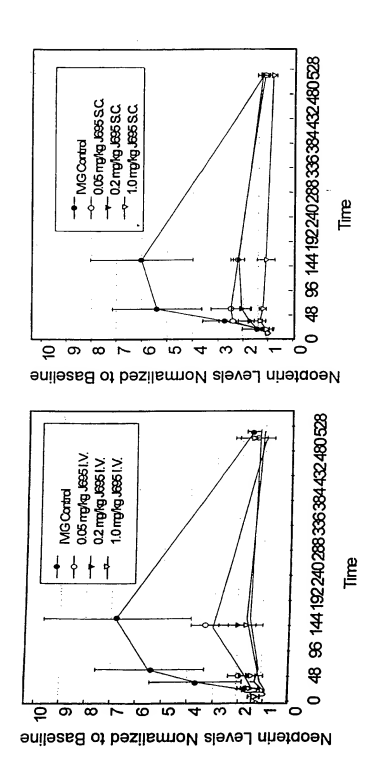


Figure 3: In vivo efficacy of J695 in cynomolgus monkeys



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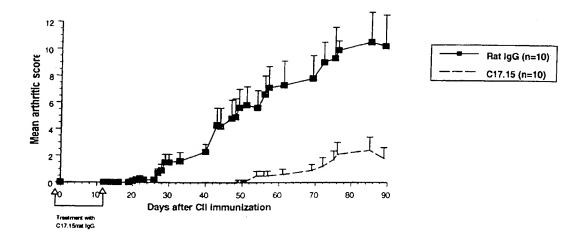


FIGURE 4

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<170> PatentIn Ver. 2.0
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<210> 2
<211> 12
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<223> Xaa at position 9 could be either Ser or Ala
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<223> Xaa at position 10 could be either Asn, Gly or Tyr
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<223> Xaa at position 11 could be either Thr or Asp
<223> Xaa at position 13 could be either Lys or His
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<223> Xaa at position 16 could be either Arg or Gly
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<223> Xaa at position 31 could be either Ser or Glu
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<223> Xaa at position 84 could be either Lys or Asn
<223> Xaa at position 97 could be either Thr, Ala or Lys
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<223> Xaa at position 98 could be either Thr or Lys
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                                     10
```

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Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ala Phe Ile Arg Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Asx 50 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Xaa Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys $85 \hspace{1cm} 90 \hspace{1cm} 95$

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Val Ser Ser 115

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- 5 -

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Xaa Xaa Xaa Xaa Phe Gly Thr Gly Thr Lys Val Thr Val Leu Gly 105 100

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<223> Xaa at position 3 could be either Ser or Thr
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<223> Xaa at position 5 could be either Asp or Ser
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His Xaa Xaa Xaa Xaa
 1
<210> 10
<211> 12
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     Leu, Phe, Arg, His, Asn or Tyr
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Gln Ser Tyr Xaa Xaa Xaa Thr His Pro Ala Leu Leu
                 5
<210> 11
<211> 17
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<223> Xaa at position 5 could be either Asp, Ser, Glu or
     Ala
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<223> Xaa at position 6 could be either Gly or Arg
<223> Xaa at position 8 represents any amino acid
```

```
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<400> 11
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                                    10
                  5
Gly
<210> 12
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<212> PRT
<213> Homo sapiens
<223> Xaa at position 1 could be either Gly, Tyr, Ser,
    Thr, Asn or Gln
<400> 12
Xaa Asn Asp Gln Arg Pro Ser
<210> 13
<211> 9
<212> PRT
<213> Homo sapiens
<223> Xaa at position 4 and 5 represents any amino acid
<223> Xaa at position 6 could be either Tyr or His
<223> Xaa at position 7 could be either Gly, Met, Ala,
     Asn or Ser
<400> 13
Phe Thr Phe Xaa Xaa Xaa Met His
         5
<210> 14
<211> 13
<212> PRT
<213> Homo sapiens
<223> Xaa at position 9 could be either Ser, Cys, Arg,
     Asn, Asp or Thr
<223> Xaa at position 10 could be either Asn, Met or Ile
<220>
<223> Xaa at position 11 could be either Thr, Tyr, Asp,
     His, Lys or Pro
<400> 14
Ser Gly Gly Arg Ser Asn Ile Gly Xaa Xaa Xaa Val Lys
 1
                                    10
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<210> 15

<211> 114

<212> PRT

<213> Homo sapiens

<220>

<223> Xaa at position 30 could be Ser or Glu

<220>

<223> Xaa at position 83 could be Lys or Asn

<223> Xaa at position 5 could be either Gln or Glu

<400> 15

Gln Val Gln Val Xaa Ser Gly Gly Gly Val Val Gln Pro Gly Arg Ser

Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Xaa Tyr Gly

Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val Ala

Phe Ile Arg Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val Lys 55

Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr Leu 70

Gln Met Xaa Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys Lys

Thr His Gly Ser His Asp Asn Trp Gly Gln Gly Thr Met Val Thr Val 105

Ser Ser

<210> 16

<211> 112

<212> PRT

<213> Homo sapiens

<223> Xaa at position 1 could be either Ser or Gln

<223> Xaa at position 2 could be Tyr or Ser

<220>

<223> Xaa at position 13 could be either Thr or Ala

<223> Xaa at position 25 could be either Gly or Ser

<223> Xaa at position 51 and 95 could be either Gly or Tyr

<223> Xaa at position 79 could be either Val or Leu

<400> 16

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Xaa Xaa Val leu Thr Gln Pro Pro Ser Val Ser Gly Xaa Pro Gly Gln
Arg Val Thr Ile Ser Cys Ser Gly Xaa Arg Ser Asn Ile Gly Ser Asn
             20
                                  25
Thr Val Lys Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu
Ile Tyr Xaa Asn Asp Gln Arg Pro Ser Gly Val Pro Asp Arg Phe Ser
Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Thr Gly Xaa Gln
                                         75
Ala Glu Asp Glu Ala Asp Tyr Tyr Cys Gln Ser Tyr Asp Arg Xaa Thr
His Pro Ala Leu Leu Phe Gly Thr Gly Thr Lys Val Thr Val Leu Gly
<210> 17
<211> 6
<212> PRT
<213> Homo sapiens
<400> 17
His Gly Ser His Asp Asn
 1
<210> 18
<211> 12
<212> PRT
<213> Homo sapiens
<400> 1.8
Gln Ser Tyr Asp Arg Gly Thr His Pro Ala Leu Leu
<210> 19
<211> 17
<212> PRT
<213> Homo sapiens
<400> 19
Phe Ile Arg Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val Lys
Gly
<210> 20
<211> 7
<212> PRT
<213> Homo sapiens
<400> 20
Gly Asn Asp Gln Arg Pro Ser
```

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<210> 21
<211> 9
<212> PRT
<213> Homo sapiens
<400> 21
Phe Thr Phe Ser Ser Tyr Gly Met His
<210> 22
<211> 13
<212> PRT
<213> Homo sapiens
<400> 22
Ser Gly Gly Arg Ser Asn Ile Gly Ser Asn Thr Val Lys
                                    1.0
<210> 23
<211> 115
<212> PRT
<213> Homo sapiens
<400> 23
Gln Val Gln Leu Val Gln Ser Gly Gly Gly Val Val Gln Pro Gly Arg
                                  10
Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr
Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val
                            40
Ala Phe Ile Arg Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val
Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr
Leu Gln Met Lys Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys
Lys Thr His Gly Ser His Asp Asn Trp Gly Gln Gly Thr Met Val Thr
                               105
Val Ser Ser
      115
<210> 24
<211> 112
<212> PRT
<213> Homo sapiens
Ser Tyr Val Leu Thr Gln Pro Pro Ser Val Ser Gly Thr Pro Gly Gln
Arg Val, Thr Ile Ser Cys Ser Gly Gly Arg Ser Trp Ile Gly Ser Asn
Thr Val Lys Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu
```

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35 45 Ile Tyr Gly Asn Asp Gln Arg Pro Ser Gly Val Pro Asp Arg Phe Ser Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Thr Gly Val Gln Ala Glu Asp Glu Ala Asp Tyr Tyr Cys Gln Ser Tyr Asp Arg Gly Thr 85 His Pro Ala Leu Leu Phe Gly Thr Gly Thr Lys Val Thr Val Leu Gly <210> 25 <211> 6 <212> PRT <213> Homo sapiens <400> 25 His Gly Ser His Asp Asn 1 5 <210> 26 <211> 12 <212> PRT <213> Homo sapiens <400> 26 Gln Ser Tyr Asp Arg Tyr Thr His Pro Ala Leu Leu <210> 27 <211> 17 <212> PRT <213> Homo sapiens Phe Ile Arg Tyr Asp Gly Ser Asm Lys Tyr Tyr Ala Asp Ser Val Lys Gly <210> 28 <211> 7 <212> PRT <213> Homo sapiens <400> 28 Tyr Asn Asp Gln Arg Pro Ser <210> 29 <211> 9 <212> PRT <213> Homo sapiens

```
<400> 29
  Phe Thr Phe Ser Ser Tyr Gly Met His
  <210> 30
  <211> 13
  <212> PRT
  <213> Homo sapiens
  <400> 30
  Ser Gly Ser Arg Ser Asn Ile Gly Ser Asn Thr Val Lys
  <210> 31
 <211> 115
 <212> PRT
 <213> Homo sapiens
 <400> 31
 Gln Val Gln Leu Val Glu Ser Gly Gly Val Val Gln Pro Gly Arg
                                      10
 Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr
 Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val
 Ala Phe Ile Arg Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val
 Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr
 Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys
Lys Thr His Gly Ser His Asp Asn Trp Gly Gln Gly Thr Met Val Thr
                               105
 Val Ser Ser
        115
<210> 32
<211> 112
<212> PRT
<213> Homo sapiens
<400> 32
Gln Ser Val Leu Thr Gln Pro Pro Ser Val Ser Gly Ala Pro Gly Gln
Arg Val Thr Ile Ser Cys Ser Gly Ser Arg Ser Asn Ile Gly Ser Asn
                                25
Thr Val Lys Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu
Ile Tyr Tyr Asn Asp Gln Arg Pro Ser Gly Val Pro Asp Arg Phe Ser
Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Thr Gly Leu Gln
                    70
```

Ala Glu Asp Glu Ala Asp Tyr Tyr Cys Gln Ser Tyr Asp Arg Tyr Thr 85 90 95

His Pro Ala Leu Leu Phe Gly Thr Gly Thr Lys Val Thr Val Leu Gly 100 105 110

<210> 33

<211> 115

<212> PRT

<213> Homo sapiens

<400> 33

Gln Val Gln Leu Val Gln Ser Gly Gly Gly Val Val Gln Pro Gly Arg 1 5 10

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ala Phe Ile Arg Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 55 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Lys Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys 85 90 95

Thr Thr Ser Gly Ser Tyr Asp Tyr Trp Gly Gln Gly Thr Met Val Thr 100 105 110

Val Ser Ser 115

<210> 34

<211> 112

<212> PRT

<213> Homo sapiens

<400> 34

Ser Tyr Val Leu Thr Gln Pro Pro Ser Val Ser Gly Thr Pro Gly Gln 1 5 10

Arg Val Thr Ile Ser Cys Ser Gly Gly Arg Ser Asn Ile Gly Ser Asn 20 25 30

Thr Val Lys Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu 35 40 45

Ile Tyr Gly Asn Asp Gln Arg Pro Ser Gly Val Pro Asp Arg Phe Ser 50 55 60

Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Thr Gly Val Gin 65 70 75 80

Ala Glu Asp Glu Ala Asp Tyr Tyr Cys Gln Ser Tyr Asp Ser Ser Leu 85 90 95

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Arg Gly Ser Arg Val Phe Gly Thr Gly Thr Lys Val Thr Val Leu Gly 105

<210> 35

<211> 115

<212> PRT

<213> Homo sapiens

<400> 35

Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Gly

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val

Ala Phe Ile Arg Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys

Ala Lys Ser Gly Ser Tyr Asp Tyr Trp Gly Gln Gly Thr Met Val Thr

Val Ser Ser 115

<210> 36

<211> 112

<212> PRT

<213> Homo sapiens

<223> Xaa at position 32 represents either Gly or Tyr

<400> 36

Gln Ser Val Leu Thr Gln Pro Pro Ser Val Ser Gly Ala Pro Gly Gln

Arg Val Thr Ile Ser Cys Thr Gly Ser Ser Ser Asn Ile Gly Ala Xaa

Asp Val His Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu

Ile Tyr Gly Asn Ser Asn Arg Pro Ser Gly Val Pro Asp Arg Phe Ser

Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Thr Gly Leu Gln

Ala Glu Asp Glu Ala Asp Tyr Tyr Cys Gln Ser Tyr Asp Ser Ser Leu

Ser Gly Ser Arg Val Phe Gly Thr Gly Thr Lys Val Thr Val Leu Gly

110

100 105 .

<210> 37 <211> 115 <212> PRT

<213> Homo sapiens

<400> 37

Gln Val Gln Leu Val Gln Ser Gly Gly Gly Val Val Gln Pro Gly Arg 1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr \$20\$ \$25\$ \$30\$ \cdot

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ala Phe Iie Arg Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Lys Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cỳs $85 \hspace{1cm} 90 \hspace{1cm} 95$

Thr Thr His Gly Ser His Asp Asr Trp Gly Gln Gly Thr Met Val Thr $100 \hspace{1.5cm} 105 \hspace{1.5cm} 110$

Val Ser Ser 115

<210> 38

<211> 112 <212> PRT

<213> Homo sapiens

<400> 38

Ser Tyr Val Leu Thr Gln Pro Pro Ser Val Ser Gly Thr Pro Gly Gln 1 5 10 15

Arg Val Thr Ile Ser Cys Ser Gly Gly Arg Ser Asn Ile Gly Ser Asn 20 25 30

Thr Val Lys Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu 35 40 45

Ile Tyr Gly Asn Asp Gln Arg Pro Ser Gly Val Pro Asp Arg Phe Ser 50 55 60

Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Thr Gly Val Gln 65 70 75 80

Ala Glu Asp Glu Ala Asp Tyr Tyr Cys Gln Ser Tyr Asp Ser Ser Leu 85 . 90 95

Arg Gly Ser Arg Val Phe Gly Thr Gly Thr Lys Val Thr Val Leu Gly $100 \hspace{1cm} 105 \hspace{1cm} 110$

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<210> 39 <211> 115 <212> PRT <213> Homo sapiens

<400> 39

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Gly Met His Trp Val Arg Gln Aia Pro Gly Lys Gly Leu Glu Trp Val 35 46 45

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Lys Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys $85 \hspace{1cm} 90 \hspace{1cm} 95$

Thr Thr Ser Gly Ser Tyr Asp Tyr Trp Gly Gln Gly Thr Met Val Thr 100 105 110

Val Ser Ser 115

<210> 40

<211> 112

<212> PRT

<213> Homo sapiens

<400> 40

Ser Tyr Val Leu Thr Gln Pro Prc Ser Val Ser Gly Thr Pro Gly Gln $1 \hspace{1cm} 5 \hspace{1cm} 10 \hspace{1cm} 15$

Arg Val Thr Ile Ser Cys Ser Gly Gly Arg Ser Asn Ile Gly Ser Asn 20 25 30

Thr Val Lys Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu 35 40 45

Ile Tyr Gly Asn Asp Gln Arg Pro Ser Gly Val Pro Asp Arg Phe Ser 50 60

Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Thr Gly Val Gln 65 70 75 80

Ala Glu Asp Glu Ala Asp Tyr Tyr Cys Gln Ser Tyr Asp Arg Gly Phe
85 90 95

Thr Gly Ser Arg Val Phe Gly Thr Gly Thr Lys Val Thr Val Leu Gly 100 105 110

<210> 41 <211> 115

- 17 -

<212> PRT <213> Homo sapiens

<400> 41

Gln Val Gln Leu Val Gln Ser Gly Gly Gly Val Val Gln Pro Gly Arg 1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ala Phé Ile Arg Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 55 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Lys Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys 85 90 95

Thr Thr Ser Gly Ser Tyr Asp Tyr Trp Gly Gln Gly Thr Met Val Thr 100 105 110

Val Ser Ser 115

<210> 42

<211> 112

<212> PRT

<213> Homo sapiens

<400> 42

Ser Tyr Val Leu Thr Gln Pro Pro Ser Val Ser Gly Thr Pro Gly Gln
1 10 15

Arg Val Thr Ile Ser Cys Ser Gly Gly Arg Ser Asn Ile Gly Ser Asn 20 25 30

Thr Val Lys Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu
35 40

Ile Tyr Gly Asn Asp Gln Arg Pro Ser Gly Val Pro Asp Arg Phe Ser

Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Thr Gly Val Gln
65 70 75

Ala Glu Asp Glu Ala Asp Tyr Tyr Cys Gln Ser Tyr Asp Ser Ser Leu 85 90 . 95

Trp Gly Ser Arg Val Phe Gly Thr Gly Thr Lys Val Thr Val Leu Gly 100 105 110

<210> 43

<211> 115

<212> PRT

<213> Homo sapiens

<400> 43

Gln Val Gln Leu Val Gln Ser Gly Gly Gly Val Val Gln Pro Gly Arg 1 5 10

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val
35 40 45

Ala Phe Ile Arg Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 55 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Lys Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys 85 90 95

Thr Thr His Gly Ser His Asp Asn Trp Gly Gln Gly Thr Met Val Thr 100 105 110

Val Ser Ser 115

<210> 44

<211> 112

<212> PRT

<213> Homo sapiens

<400> 44

Ser Tyr Val Leu Thr Gln Pro Pro Ser Val Ser Gly Thr Pro Gly Gln 1 5 10

Arg Val Thr Ile Ser Cys Ser Gly Gly Arg Ser Asn Ile Gly Ser Asn 20 25 30

Thr Val Lys Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu $35 \hspace{1cm} 40 \hspace{1cm} 45$

Ile Tyr Gly Asn Asp Gln Arg Pro Ser Gly Val Pro Asp Arg Phe Ser
50 55 60

Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Thr Gly Val Gln 65 70 75 80

Ala Glu Asp Glu Ala Asp Tyr Tyr Cys Gln Ser Tyr Asp Arg Gly Phe
85 90 95

Thr Gly Ser Arg Val Phe Gly Thr Gly Thr Lys Val Thr Val Leu Gly 100 105 110

<210> 45

<211> 115

<212> PRT

<213> Homo sapiens

<400> 45

Gin Val Gln Leu Val Gln Ser Gly Gly Gly Val Val Gln Pro Gly Arg

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr

20 25 30

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ala Phe Ile Arg Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 55 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 . 70 75 80

Leu Gln Met Lys Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys 85 90 95

Thr Thr His Gly Ser His Asp Asn Trp Gly Gln Gly Thr Met Val Thr 100 105 110

Val Ser Ser 115

<210> 46

<211> 112

<212> PRT

<213> Homo sapiens

<400> 46

Ser Tyr Val Leu Thr Gln Pro Pro Ser Val Ser Gly Thr Pro Gly Gln 1 5 10 15

Arg Val Thr Ile Ser Cys Ser Gly Gly Arg Ser Asn Ile Gly Ser Asn 20 25 30

Thr Val Lys Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu 35 40 45

Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Thr Gly Val Gln
65 70 75 80

Ala Glu Asp Glu Ala Asp Tyr Tyr Cys Gln Ser Tyr Asp Ser Ser Leu 85 90 95

Trp Gly Ser Arg Val Phe Gly Thr Gly Thr Lys Val Thr Val Leu Gly 100 105 \cdot 110

<210> 47

<211> 115

<212> PRT

<213> Homo sapiens

<400> 47

Gln Val Gln Leu Val Gln Ser Gly Gly Gly Val Val Gln Pro Gly Arg 1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr \$20\$ \$25\$ \$30

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35, 40 45

Ala Phe Ile Arg Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 .75 80

Leu Gln Met Lys Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys
85 90 95

Lys Thr His Gly Ser His Asp Asn Trp Gly Gln Gly Thr Met Val Thr 100 105 110

Val Ser Ser 115

<210> 48

<211> 112

<212> PRT

<213> Homo sapiens

<400> 48

Ser Tyr Val Leu Thr Gln Pro Pro Ser Val Ser Gly Thr Pro Gly Gln 1 5 10 15

Arg Val Thr Ile Ser Cys Ser Gly Ser Arg Ser Asn Ile Gly Ser Asn 20 25 30

Thr Val Lys Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu 35 40 45

Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Thr Gly Val Gln 65 70 75 80

Ala Glu Asp Glu Ala Asp Tyr Tyr Cys Gln Thr Tyr Asp Lys Gly Phe
85 90 95

Thr Gly Ser Ser Val Phe Gly Thr Gly Thr Lys Val Thr Val Leu Gly
100 105 110

<210> 49

<211> 115

<212> PRT

<213> Homo sapiens

<400> 49

Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg 1 5 10

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val

Ala Phe Ile Arg Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val $50 \hspace{1cm} 55 \hspace{1cm} 60 \hspace{1cm}$

- 21 -

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 70

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys

Lys Thr His Gly Ser His Asp Asn Trp Gly Gln Gly Thr Met Val Thr 105

Val Ser Ser 115

<210> 50

<211> 112

<212> PRT

<213> Homo sapiens

<400> 50

Gln Ser Val Leu Thr Gln Pro Pro Ser Val Ser Gly Ala Pro Gly Gln

Arg Val Thr Ile Ser Cys Ser Gly Ser Arg Ser Asn Ile Gly Ser Asn

Thr Val Lys Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu

Ile Tyr Gly Asn Asp Gln Arg Pro Ser Gly Val Pro Asp Arg Phe Ser

Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Thr Gly Leu Gln

Ala Glu Asp Glu Ala Asp Tyr Tyr Cys Gln Thr Tyr Asp Lys Gly Phe

Thr Gly Ser Ser Val Phe Gly Thr Gly Thr Lys Val Thr Val Leu Gly 105

<210> 51

<211> 115

<212> PRT

<213> Homo sapiens

<400> 51

Gln Val Gln Leu Val Gln Ser Gly Gly Gly Val Val Gln Pro Gly Arg

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val

Ala Phe Ile Arg Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr

Leu Gln Met Lys Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys

- 22 -

85 90 95

Thr Thr His Gly Ser His Asp Thr Trp Gly Gln Gly Thr Met Val Thr 100 105 110

Val Ser Ser 115

<210> 52

<211> 112

<212> PRT

<213> Homo sapiens

<400> 52

Ser Tyr Val Leu Thr Gln Pro Pro Ser Val Ser Gly Thr Pro Gly Gln $1 \hspace{1cm} 5 \hspace{1cm} 10 \hspace{1cm} 15$

Arg Val Thr Ile Ser Cys Ser Gly Gly Arg Ser Asn Ile Gly Ser Asn 20 25 30

Thr Val Lys Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu 35 40 45

Ile Tyr Gly Asn Asp Gln Arg Pro Ser Gly Val Pro Asp Arg Phe Ser 50 60

Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Thr Gly Val Gln 65 70 75 80

Ala Glu Asp Glu Ala Asp Tyr Tyr Cys Gln Ser Tyr Asp Ser Ser Leu 85 90 95

Trp Gly Thr Arg Val Phe Gly Thr Gly Thr Lys Val Thr Val Leu Gly 100 105 110

<210> 53

<211> 115

<212> PRT

<213> Homo sapiens

<400> 53

Gln Val Gln Leu Val Gln Ser Gly Gly Gly Val Val Gln Pro Gly Arg 1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ala Phe Ile Arg Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 55 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Lys Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys 85 90 95

Thr Thr His Gly Ser His Asp Asn Trp Gly Gln Gly Thr Met Val Thr 100 105 110 - 23 -

Val Ser Ser 115

<210> 54

<211> 112

<212> PRT

<213> Homo sapiens

<400> 54

Ser Tyr Val Leu Thr Gln Pro Pro Ser Val Ser Gly Thr Pro Gly Gln

Arg Val Thr Ile Ser Cys Ser Gly Gly Arg Ser Asn Ile Val Ser Asn

Thr Val Lys Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu

Ile Tyr Gly Asn Asp Gln Arg Pro Ser Gly Val Pro Asp Arg Phe Ser

Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Thr Gly Val Gln

Ala Glu Asp Glu Ala Asp Tyr Tyr Cys Gln Ser Tyr Asp Arg Gly Phe

Thr Gly Ser Arg Val Phe Gly Thr Gly Thr Lys Val Thr Val Leu Gly 105

<210> 55

<211> 115

<212> PRT

<213> Homo sapiens

<400> 55

Gln Val Gln Leu Val Gln Ser Gly Gly Gly Val Val Gln Pro Gly Arg

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val

Ala Phe Ile Arg Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr

Leu Gln Met Lys Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys

Thr Thr His Gly Ser His Asp Asn Trp Gly Gln Gly Thr Met Val Thr

Val Ser Ser

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<210> 56

<211> 112 <212> PRT

<213> Homo sapiens

Ser Tyr Val Leu Thr Gln Pro Pro Ser Val Ser Gly Thr Pro Gly Gln

Arg Val Thr Ile Ser Cys Ser Gly Gly Arg Ser Asn Ile Val Ser Asn

Thr Val Lys Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu

Ile Tyr Gly Asn Asp Gln Arg Pro Ser Gly Val Pro Asp Arg Phe Ser

Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Thr Gly Val Gln

Ala Glu Asp Glu Ala Asp Tyr Tyr Cys Gln Ser Tyr Asp Arg Gly Phe

Thr Gly Ala Arg Val Phe Gly Thr Gly Thr Lys Val Thr Val Leu Gly 105

<210> 57

<211> 115

<212> PRT

<213> Homo sapiens

<400> 57

Gln Val Gln Leu Val Gln Ser Gly Gly Gly Val Val Gln Pro Gly Arg 10

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val

Ala Phe Ile Arg Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr

Leu Gln Met Lys Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys

Lys Thr His Gly Ser His Asp Asn Trp Gly Gln Gly Thr Met Val Thr 105

Val Ser Ser

<210> 58

<211> 112

<212> PRT

<213> Homo sapiens

<400> 58

Ser Tyr Val Leu Thr Gln Pro Pro Ser Val Ser Gly Thr Pro Gly Gln 1 5 10 15

Arg Val Thr Ile Ser Cys Ser Gly Gly Arg Ser Asn Ile Gly Ser Asn 20 25 30

Thr Val Lys Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu 35 40 45

Ile Tyr Gly Asn Asp Gln Arg Pro Ser Gly Val Pro Asp Arg Phe Ser
50 55 60

Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Thr Gly Val Gin 65 70 75 80

Ala Glu Asp Glu Ala Asp Tyr Tyr Cys Gln Thr Tyr Asp Lys-Gly Fhe 85 90 95

Thr Gly Ser Ser Val Phe Gly Thr Gly Thr Lys Val Thr Val Leu Gly 100 105 110

<210> 59

<211> 115

<212> PRT

<213> Homo sapiens

<400> 59

Gln Val Gln Leu Val Gln Ser Gly Gly Gly Val Val Gln Pro Gly Arg

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ala Phe Ile Arg Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Lys Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys

Lys Thr His Gly Ser His Asp Asn Trp Gly Gln Gly Thr Met Val Thr 100 105 110

Val Ser Ser 115

<210> 60

<211> 112

<212> PRT

<213> Homo sapiens

<400> 60

Ser Tyr Val Leu Thr Gln Pro Pro Ser Val Ser Gly Thr Pro Gly Gln

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10 15 Arg Val Thr Ile Ser Cys Ser Gly Gly Arg Ser Asn Ile Gly Ser Asn Thr Val Lys Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu 35 40 Ile Tyr Gly Asn Asp Gln Arg Pro Ser Gly Val Pro Asp Arg Phe Ser Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Thr Gly Val Gln Ala Glu Asp Glu Ala Asp Tyr Tyr Cys Gln Ser Tyr Glu Arg Gly Phe Thr Gly Ser Met Val Phe Gly Thr Gly Thr Lys Val Thr Val Leu Gly 105 <210> 61 <211> 115 <212> PRT <213> Homo sapiens <400> 61 Gln Val Gln Leu Val Gln Ser Gly Gly Gly Val Val Gln Pro Gly Arg 10 Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val Ala Phe Ile Arg Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr Leu Gln Met Lys Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys Lys Thr His Gly Ser His Asp Asn Trp Gly Gln Gly Thr Met Val Thr 105 Val Ser Ser 115 <210> 62 <211> 112 <212> PRT <213> Homo sapiens <400> 62 Ser Tyr Val Leu Thr Gln Pro Pro Ser Val Ser Gly Thr Pro Gly Gln Arg Val Thr Ile Ser Cys Ser Gly Gly Arg Ser Asn Ile Gly Ser Asn

25

Thr Val Lys Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu 35 . 40 . 45

Ile Tyr Gly Asn Asp Gln Arg Pro Ser Gly Val Pro Asp Arg Phe Ser
50 60

Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Thr Gly Val Gin 65 70 75 80

Ala Glu Asp Glu Ala Asp Tyr Tyr Cys Gln Ser Tyr Asp Arg Gly Thr $85\,$ $\cdot\,$ 90 $95\,$

His Pro Leu Thr Ile Phe Gly Thr Gly Thr Lys Val Thr Val Leu Gly $100 \,$ $105 \,$ $110 \,$

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<211> 115

<212> PRT

<213> Homo sapiens

<400> 63

Gln Val Gln Leu Val Gln Ser Gly Gly Gly Val Val Gln Pro Gly Arg 1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr
20 25 30

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val
35 40 45

Ala Phe Ile Arg Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 55 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Lys Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys 85 90 95

Lys Thr His Gly Ser His Asp Asn Trp Gly Gln Gly Thr Met Val Thr 100 105 110

Val Ser Ser 115

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<211> 112

<212> PRT

<213> Homo sapiens

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Ser Tyr Val Leu Thr Gln Pro Pro Ser Val Ser Gly Thr Pro Gly Gln

Arg Val Thr Ile Ser Cys Ser Gly Gly Arg Ser Asn Ile Gly Ser Asn 20 25 30

Thr Val Lys Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu 35 40 45

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Ile Tyr Gly Asn Asp Gln Arg Pro Ser Gly Val Pro Asp Arg Phe Ser
50 60

Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Thr Gly Val Gln 65 70 75 80

Ala Glu Asp Glu Ala Asp Tyr Tyr Cys Gln Ser Tyr Asp Arg Gly Ser 85 90 95

His Pro Ala Leu Thr Phe Gly Thr Gly Thr Lys Val Thr Val Leu Gly $100 \,$ $105 \,$ $110 \,$

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<211> 115

<212> PRT

<213> Homo sapiens

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Gln Val Gln Leu Val Gln Ser Gly Gly Gly Val Val Gln Pro Gly Arg 1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ala Phe Ile Arg Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 55 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Lys Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys
85 90 95

Lys Thr His Gly Ser His Asp Asn Trp Gly Gln Gly Thr Met Val Thr $100 \hspace{1.5cm} 105 \hspace{1.5cm} 110$

Val Ser Ser

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<211> 112 <212> PRT

<213> Homo sapiens

<400> 66

Ser Tyr Val Leu Thr Gln Pro Pro Ser Val Ser Gly Thr Pro Gly Gln
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Arg Val Thr Ile Ser Cys Ser Gly Gly Arg Ser Asn Ile Gly Ser Asn 20 25 30

Thr Val Lys Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu 35 40 45

Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Thr Gly Val Gln

- 29 -

65 70 75 50

Ala Glu Asp Glu Ala Asp Tyr Tyr Cys Gln Ser Tyr Asp Arg Gly Thr 90

His Pro Leu Thr Met Phe Gly Thr Gly Thr Lys Val Thr Val Leu Gly 100

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Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys 85 90 95

Lys Thr His Gly Ser His Asp Asn Trp Gly Gln Gly Thr Met Val Thr $100 \\ 105 \\ 110$

Val Ser Ser 115

<210> 68 <211> 112 <212> PRT <213> Homo sapiens

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1 5 10 15

Arg Val Thr Ile Ser Cys Ser Gly Ser Arg Ser Asn Ile Gly Ser Asn 20 25 30

Thr Val Lys Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu 35 40 45

Ile Tyr Gly Asn Asp Gln Arg Pro Ser Gly Val Pro Asp Arg Phe Ser 50 60

Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Thr Gly Leu Gln 65 70 75 80

Ala Glu Asp Glu Ala Asp Tyr Tyr Cys Gln Ser Tyr Asp Arg Gly Thr 85 90 95 - 30 -

His Pro Leu Thr Met Phe Gly Thr Gly Thr Lys Val Thr Val Leu Gly 100 105

<210> 69

<211> 115

<212> PRT

<213> Homo sapiens

<400> 69

Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val

Ala Phe Ile Arg Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys

Lys Thr His Gly Ser His Asp Asn Trp Gly Gln Gly Thr Met Val Thr 105

Val Ser Ser 115

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Gln Ser Val Leu Thr Gln Pro Pro Ser Val Ser Gly Ala Pro Gly Gln

Arg Val Thr Ile Ser Cys Ser Gly Ser Arg Ser Asn Ile Gly Ser Asn

Thr Val Lys Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu

Ile Tyr Gly Asn Asp Gln Arg Pro Ser Gly Val Pro Asp Arg Phe Ser

Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Thr Gly Leu Gln

Ala Glu Asp Glu Ala Asp Tyr Tyr Cys Gln Ser Tyr Asp Arg Gly Thr 90

His Pro Ala Leu Leu Phe Gly Thr Gly Thr Lys Val Thr Val Leu Gly 105

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Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Glu Tyr
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Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val
Ala Phe Ile Arg Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val
Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr
Leu Gln Met Lys Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys
Lys Thr His Gly Ser His Asp Asn Trp Gly Gln Gly Thr Met Val Thr
                     105
Val Ser Ser
      115
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Gln Ser Val Leu Thr Gln Pro Pro Ser Val Ser Gly Ala Pro Gly Gln
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Arg Val Thr Ile Ser Cys Ser Gly Ser Arg Ser Asn Ile Gly Ser Asn
Thr Val Lys Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu
Ile Tyr Gly Asn Asp Gln Arg Pro Ser Gly Val Pro Asp Arg Phe Ser
Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Thr Gly Leu Gln
Ala Glu Asp Glu Ala Asp Tyr Tyr Cys Gln Ser Tyr Asp Arg Gly Thr
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His Pro Ala Leu Leu Phe Gly Thr Gly Thr Lys Val Thr Val Leu Gly

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<211> 115

<212> PRT

<213> Homo sapiens

<400> 73

Gln Val Gln Leu Val Gln Ser Gly Gly Gly Val Val Gln Pro Gly Arg 1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ala Phe Ile Arg Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val50 $\,$ 60 $\,\cdot$

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Lys Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys 85 90 95

Lys Thr His Gly Ser His Asp Asn Trp Gly Gln Gly Thr Met Val Thr $100 \hspace{1cm} 105 \hspace{1cm} 110$

Val Ser Ser 115

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<212> PRT

<213> Homo sapiens

<400> 74

Gln Ser Val Leu Thr Gln Pro Pro Ser Val Ser Gly Ala Pro Gly Gln 1 5 10 15

Arg Val Thr Ile Ser Cys Ser Gly Ser Arg Ser Asn Ile Gly Ser Asn 20 25 30

Thr Val Lys Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu 35 40 45

Ile Tyr Tyr Asn Asp Gln Arg Pro Ser Gly Val Pro Asp Arg Phe Ser 50 60

Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Thr Gly Leu Gln
65 70 75 80

Ala Glu Asp Glu Ala Asp Tyr Tyr Cys Gln Ser Tyr Asp Arg Gly Thr
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His Pro Ala Leu Leu Phe Gly Thr Gly Thr Lys Val Thr Val Leu Gly 100 105 110

<210> 75

<211> 115

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- 33 -<400> 75 Gln Val Gln Leu Val Gln Ser Gly Gly Gly Val Val Gln Pro Gly Arg Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val Ala Phe Ile Arg Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 70 Leu Gln Met Lys Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys Lys Thr His Gly Ser His Asp Asn Trp Gly Gln Gly Thr Met Val Thr Val Ser Ser 115 <210> 76 <211> 112 <212> PRT <213> Homo sapiens <400> 76 Gln Ser Val Leu Thr Gln Pro Pro Ser Val Ser Gly Ala Pro Gly Gln

Arg Val Thr Ile Ser Cys Ser Gly Ser Arg Ser Asn Ile Gly Ser Asn

Thr Val Lys Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu 40

Ile Tyr Gly Asn Asp Gln Arg Pro Ser Gly Val Pro Asp Arg Phe Ser

Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Thr Gly Leu Gln

Ala Glu Asp Glu Ala Asp Tyr Tyr Cys Gln Ser Tyr Asp Arg Tyr Thr

His Pro Ala Leu Leu Phe Gly Thr Gly Thr Lys Val Thr Val Leu Gly 105

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His Gly Ser Tyr Asp Tyr
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Arg Arg Arg Ser Asn Tyr
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His Gly Ser His Asp Asp
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His Gly Ser His Asp Asn
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Ala Lys His Gly Ser His Asp Asn Trp Gly Gln Gly
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Thr Thr His Gly Ser His Asp Asr. Trp Ser Gln Gly
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Lys Thr His Gly Ser His Asp Asn Trp Gly Gln Gly
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Lys Thr His Gly Ser His Asp Asn Trp Gly His Gly
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Thr Thr His Gly Ser His Asp Asn Trp Ser Gln Gly
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Thr Thr His Gly Ser His Asp Thr
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Thr Lys His Gly Ser His Asp Asn
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Thr Thr Gln Gly Arg His Asp Asn
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Lys Thr Arg Gly Arg His Asp Asn
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 Thr Thr His Gly Ser His Asp Asp
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<211> 8
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Lys Thr His Gly Ser His Asp Asn
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Lys Thr His Gly Ser His Asp Asn
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Thr Thr His Gly Ser His Asp Asn
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Ala Thr His Gly Ser Gln Asp Asn
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His Gly Ser Gln Asp Thr
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Ser Gly Ser Tyr Asp Tyr
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His Gly Ser Gln Asp Asn
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Cys Lys Thr His Gly Ser His Asp Asn
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Gln Ser Tyr Asp Ser Ser Leu Trp Gly Thr Arg Val
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Gln Ser Tyr Asp Arg Asp Phe Thr Gly Ser Arg Val
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Gln Ser Tyr Asp Arg Arg Phe Thr Gly Ser Arg Val
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His Gly Ser His Asp Phe
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Cys Asn Asp Gln Arg Pro Ser
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Ser Asn Asp Gln Arg Pro Ser
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Ala Asn Asp Gln Arg Pro Ser
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Gly Asn Asp Ser Arg Pro Ser
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Gln Ser Tyr Cys Arg Gly Thr His Pro Ala Leu Leu
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Gln Ser Tyr Ser Arg Gly Thr His Pro Ala Leu Leu
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Gln Ser Tyr Tyr Arg Gly Thr His Pro Ala Leu Leu
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Gln Ser Tyr Asn Arg Gly Thr His Pro Ala Leu Leu
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Gln Ser Tyr Ile Arg Gly Thr His Pro Ala Leu Leu
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Gln Ser Tyr Asp Asp Gly Thr His Pro Ala Leu Leu
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Gln Ser Tyr Asp Cys Gly Thr His Pro Ala Leu Leu
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Gln Ser Tyr Asp Ser Gly Thr His Pro Ala Leu Leu
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Gln Ser Tyr Asp Tyr Gly Thr His Pro Ala Leu Leu
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<213> Homo sapiens
<400> 551
Gln Ser Tyr Asp Arg Gly Thr His Pro Ala Leu Leu
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<213> Homo sapiens
Gln Ser Tyr Asp Asn Gly Thr His Pro Ala Leu Leu
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<210> 553
<211> 12
<212> PRT
<213> Homo sapiens
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\operatorname{Gln} Ser Tyr Asp \operatorname{Gln} Gly Thr His \operatorname{\tt Pro} Ala Leu Leu
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Gln Ser Tyr Asp Thr Gly Thr His Pro Ala Leu Leu
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            5
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{\tt Gln} Ser Tyr Asp Gly Gly Thr His Pro Ala Leu Leu
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<213> Homo sapiens
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Gln Ser Tyr Asp Ala Gly Thr His Pro Ala Leu Leu
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<213> Homo sapiens
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Gln Ser Tyr Asp Val Gly Thr His Pro Ala Leu Leu
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<210> 558
<211> 12
<212> PRT
<213> Homo sapiens
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Gln Ser Tyr Asp Leu Gly Thr His Pro Ala Leu Leu
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Gln Ser Tyr Asp Ile Gly Thr His Pro Ala Leu Leu
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Gln Ser Tyr Asp Pro Gly Thr His Pro Ala Leu Leu
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<213> Homo sapiens
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Gln Ser Tyr Asp Trp Gly Thr His Pro Ala Leu Leu
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Gln Ser Tyr Asp Arg Asp Thr His Pro Ala Leu Leu
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<211> 12
<212> PRT
<213> Homo sapiens
Gln Ser Tyr Asp Arg Cys Thr His Pro Ala Leu Leu
<210> 565
<211> 12
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<212> PRT
<213> Homo sapiens
Gln Ser Tyr Asp Arg Ser Thr His Pro Ala Leu Leu
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<213> Homo sapiens
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Gln Ser Tyr Asp Arg Tyr Thr His Pro Ala Leu Leu
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Gln Ser Tyr Asp Arg His Thr His Pro Ala Leu Leu
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Gln Ser Tyr Asp Arg Arg Thr His Pro Ala Leu Leu
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Gln Ser Tyr Asp Arg Asn Thr His Pro Ala Leu Leu
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Gln Ser Tyr Asp Arg Gln Thr His Pro Ala Leu Leu
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Gln Ser Tyr Asp Arg Thr Thr His Pro Ala Leu Leu
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Gln Ser Tyr Asp Arg Gly Thr His Pro Ala Leu Leu
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Gln Ser Tyr Asp Arg Ala Thr His Pro Ala Leu Leu
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Gln Ser Tyr Asp Arg Val Thr His Pro Ala Leu Leu
<210> 575
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<212> PRT
<213> Homo sapiens
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Gln Ser Tyr Asp Arg Leu Thr His Pro Ala Leu Leu
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<211> 12
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<213>.Homo sapiens
<400> 576
Gln Ser Tyr Asp Arg Ile Thr His Pro Ala Leu Leu
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Gln Ser Tyr Asp Arg Pro Thr His Pro Ala Leu Leu
<210> 578
<211> 12
<212> PRT
<213> Homo sapiens
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Gln Ser Tyr Asp Arg Trp Thr His Pro Ala Leu Leu
<210> 579
<211> 12
<212> PRT
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<400> 579
Gln Ser Tyr Asp Arg Phe Thr His Pro Ala Leu Leu
<210> 580
<211> 48
<212> DNA
<213> synthetic construct
<223> nucleotides at positions 16 to 34 can be
     substituted with any nucleotide such that the
      randomized nucleotides represent 12% of the
     sequence
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tgtcccttgg ccccagtagt catageteee actggtegta cagtaata
                                                                   48
<210> 581
<211> 35
<212> DNA
<213> synthetic construct
<400> 581
gacacctcga tcagcggata acaatttcac acagg
                                                                   35
<210> 582
<211> 15
<212> DNA
<213> synthetic construct
<400> 582
                                                                   15
tggggccaag ggaca
<210> 583
<211> 45
<212> DNA
<213> synthetic construct
attogtocta taccgttcta ctttgtcgtc tttccagacg ttagt
                                                                   45
<210> 584
<211> 18
<212> DNA
<213> synthetic construct
<400> 584
                                                                   18
attogtocta tacogtto
<210> 585
<211> 66
<212> DNA
<213> synthetic construct
<223> nucleotides from position 28 to 42 can be
     substituted with any nucleotide such that the
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randomized nucleotides represent 12% of the
      sequence
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ggtcccagtt ccgaagaccc tcgaacccct caggetgetg tcatatgact ggcagtaata 60
gtcagc
<210> 586
<211> 15
<212> DNA
<213> synthetic construct
<400> 586
                                                                    15
tggggccaag ggaca
<210> 587
<211> 24
<212> DNA
<213> synthetic construct
<400> 587
tgaagagacg gtgaccattg tccc
                                                                    24
<210> 588
<211> 16
<212> DNA
<213> synthetic construct
<400> 588
                                                                    16
gacacctcga tcagcg
<210> 589
<211> 48
<212> DNA
<213> synthetic construct
<400> 589
                                                                    48
gagtcattct cgacttgcgg ccgcacctag gacggtcagc ttggtccc
<210> 590
<211> 12
<212> PRT
<213> Homo sapiens
<400> 590
Gln Ser Tyr Asp Arg Gly Phe Thr Gly Ser Met Val
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<210> 591
<211> 12
<212> PRT
<213> Homo sapiens
<223> Xaa is encoded by a randomized codon of sequence
      NNS with N being any nucleotide and S being either
      deoxycytosine or deoxyguanidine
<400> 591
Xaa Xaa Xaa Xaa Xaa Phe Thr Gly Ser Met Val
                  5
 1
<210> 592
<211> 12
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<212> PRT
<213> Homo sapiens
<223> Xaa is encoded by a randomized codon of sequence
      NNS with N being any nucleotide and S being either
      deoxycytosine or deoxyguanidine
<400> 592
Gln Ser Tyr Xaa Xaa Xaa Xaa Xaa Xaa Ser Met Val
                  5
<210> 593
<211> 12
<212> PRT
<213> Homo sapiens
<223> Xaa is encoded by a randomized codon of sequence
      NNS with N being any nuclectide and S being either
      deoxycytosine or deoxyguanidine
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Gln Ser Tyr Asp Arg Gly Xaa Xaa Xaa Xaa Xaa Xaa
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<211> 100
<212> PRT
<213> Homo sapiens
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Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Gly
Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Asp His
Tyr Met Asp Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val
Gly Arg Thr Arg Asn Lys Ala Asn Ser Tyr Thr Thr Glu Tyr Ala Ala
Ser Val Lys Gly Arg Phe Thr Ile Ser Arg Asp Asp Ser Lys Asn Ser
Leu Tyr Leu Gln Met Asn Ser Leu Lys Thr Glu Asp Thr Ala Val Tyr
Tyr Cys Ala Arg
           100
<210> 595
<211> 100
<212> PRT
<213> Homo sapiens
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Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Gly
Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Asp His
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20 25 30

Tyr Met Ser Trp Val Arg Gln Ala Gln Gly Lys Gly Leu Glu Leu Val 35 40 45

Gly Leu Ile Arg Asn Lys Ala Asn Ser Tyr Thr Thr Glu Tyr Ala Ala 50 55 60

Ser Val Lys Gly Arg Leu Thr Ile Ser Arg Glu Asp Ser Lys Asn Thr 65 70 75 50

Leu Tyr Leu Gln Met Ser Ser Leu Lys Thr Glu Asp Leu Ala Val Tyr 85 90 95

Tyr Cys Ala Arg 100

<210> 596

<211> 100

<212> PRT

<213> Homo sapiens

<400> 596

Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Gly 1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Asp His $20 \\ 25 \\ 30$

Tyr Met Ser Trp Val Arg Gln Ala Gin Gly Lys Gly Leu Glu Leu Val 35 40 45

Gly Leu Ile Arg Asn Lys Ala Asn Ser Tyr Thr Thr Glu Tyr Ala Ala 50 60

Ser Val Lys Gly Arg Leu Thr Ile Ser Arg Glu Asp Ser Lys Asn Thr 65 70 75 80

Met Tyr Leu Gln Met Ser Asn Leu Lys Thr Glu Asp Leu Ala Val Tyr 85 90 95

Tyr Cys Ala Arg

<210> 597

<211> 100

<212> PRT

<213> Homo sapiens

<400> 597

Glu Val Gln Leu Leu Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Gly 1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Asp His $20 \hspace{1cm} 25 \hspace{1cm} 30$

Tyr Met Ser Trp Val Arg Gln Ala Gln Gly Lys Gly Leu Glu Leu Val
35 40 45

Gly Leu Ile Λrg Asn Lys Ala Asn Ser Tyr Thr Thr Glu Tyr Ala Ala 50 60

Ser Val Lys Gly Arg Leu Thr Ile Ser Arg Glu Asp Ser Lys Asn Thr 65 70 75 80 - 117 -

Leu Tyr Leu Gln Met Ser Ser Leu Lys Thr Glu Asp Leu Ala Val Tyr 85 90 95

Tyr Cys Ala Arg 100

<210> 598

<211> 98

<212> PRT

<213> Homo sapiens

<400> 598

Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Arg 1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Asp Asp Tyr
20 25 30

Ala Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ser Gly Ile Ser Trp Asn Ser Gly Ser Ile Gly Tyr Ala Asp Ser Val 50 55 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ala Lys Asn Ser Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Leu Tyr Tyr Cys
85 90 95

Ala Lys

<210> 599

<211> 98

<212> PRT

<213> Homo sapiens

<400> 599

Glu Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Arg Pro Gly Gly 1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Asp Asp Tyr 20 25 30

Gly Met Ser Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val
35 40 45

Ser Gly Ile Asn Trp Asn Gly Gly Ser Thr Gly Tyr Ala Asp Ser Val 50 55 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ala Lys Asn Ser Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Leu Tyr His Cys 85 90 95

Ala Arg

<210> 600

<211> 98

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<212> PRT
<213> Homo sapiens
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<400> 600

Glu Val Gln Leu Val Glu Ser Gly Gly Val Val Val Gln Pro Gly Gly
1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Asp Asp Tyr 20 25 30

Thr Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Lys Gly Arg Pne Thr Ile Ser Arg Asp Asn Ser Lys Asn Ser Leu Tyr 65 70 75 50

Leu Gln Met Asn Ser Leu Arg Thr Glu Asp Thr Ala Leu Tyr Tyr Cys 85 90 95

Ala Lys

<210> 601

<211> 98

<212> PRT

<213> Homo sapiens

<400> 601

Gln Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Lys Pro Gly Gly 1 5 10

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Asp Tyr 20 25 30

Tyr Met Ser Trp Ile Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ser Tyr Ile Ser Ser Ser Gly Ser Thr Ile Tyr Tyr Ala Asp Ser Val $50 \hspace{1.5cm} 55 \hspace{1.5cm} 60$

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ala Lys Asn Ser Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys 85 90 95

Ala Arg

<210> 602

<211> 98

<212> PRT

<213> Homo sapiens

<400> 602

Gln Val Gln Leu Leu Glu Ser Gly Gly Giy Leu Val Lys Pro Gly Gly \cdot 1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Asp Tyr

- 119 -

Tyr Met Ser Trp Ile Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ser Tyr Ile Ser Ser Ser Ser Ser Tyr Thr Asn Tyr Ala Asp Ser Val 50 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ala Lys Asn Ser Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys
85 90 95

Ala Arg

<210> 603

<211> 100

<212> PRT

<213> Homo sapiens

<400> 603

Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Gly . 1 5 10

Ser Leu Lys Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Gly Ser 20 25 30

Ala Met His Trp Val Arg Gln Ala Ser Gly Lys Gly Leu Glu Trp Val 35 40 45

Gly Arg Ile Arg Ser Lys Ala Asn Ser Tyr Ala Thr Ala Tyr Ala Ala 50 60

Ser Val Lys Gly Arg Phe Thr Ile Ser Arg Asp Asp Ser Lys Asn Thr 65 70 75 80

Ala Tyr Leu Gln Met Asn Ser Leu Lys Thr Glu Asp Thr Ala Val Tyr

Tyr Cys Thr Arg

<210> 604

<211> 100

<212> PRT

<213> Homo sapiens

<400> 604

Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Lys Pro Gly Gly
1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Asn Ala 20 25 30

Trp Met Ser Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Gly Arg Ile Lys Ser Lys Thr Asp Gly Gly Thr Thr Asp Tyr Ala Ala 50 55 60

Pro Val Lys Gly Arg Phe Thr Ile Ser Arg Asp Asp Ser Lys Asn Thr 65 70 75 80

Leu Tyr Leu Gln Met Asn Ser Leu Lys Thr Glu Asp Thr Ala Val Tyr

- 120 -

85 90 95

Tyr Cys Thr Thr 100

<210> 605

<211> 100

<212> PRT

<213> Homo sapiens

<400> 605

Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Lys Pro Gly Gly
1 5 . 10 15

Ser Leu Arg Leu Ser Cys Ala Ala 3er Gly Phe Thr Phe Ser Asn Ala 20 25 30

Trp Met Ser Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Gly Arg Ile Glu Ser Lys Thr Asp Gly Gly Thr Thr Asp Tyr Ala Ala 50 60

Pro Val Lys Gly Arg Phe Thr Ile Ser Arg Asp Asp Ser Lys Asn Thr 65 70 75 80

Leu Tyr Leu Gln Met Asn Ser Leu Lys Thr Glu Asp Thr Ala Val Tyr 85 90 95

Tyr Cys Thr Thr 100

<210> 606

<211> 100

<212> PRT

<213> Homo sapiens

<400> 606

Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Lys Pro Gly Gly
1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Asn Ala

Trp Met Ser Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val

Gly Arg Ile Lys Ser Lys Thr Asp Gly Gly Thr Thr Asp Tyr Ala Ala

Pro Val Lys Gly Arg Phe Thr Ile Ser Arg Asp Asp Ser Lys Asn Thr 65 70 75 80

Leu Tyr Leu Gln Met Asn Ser Leu Lys Thr Glu Asp Thr Ala Val Tyr 85 90 95

Tyr Cys Thr Thr 100

<210> 607

<211> 100

<212> PRT

<213> Homo sapiens

- 121 -

Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Lys Pro Gly Gly Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Asn Ala Trp Met Ser Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val Gly Arg Ile Lys Ser Lys Thr Asp Gly Gly Thr Thr Asn Tyr Ala Ala Pro Val Lys Gly Arg Phe Thr Ile Ser Arg Asp Asp Ser Lys Asn Thr Leu Tyr Leu Gln Met Asn Ser Leu Lys Thr Glu Asp Thr Ala Val Tyr Tyr Cys Thr Thr 100 <210> 608 <211> 100 <212> PRT <213> Homo sapiens <400> 608 Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Lys Pro Gly Gly Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Asn Ala Trp Met Asn Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val Gly Arg Ile Lys Ser Lys Thr Asp Gly Gly Thr Thr Asp Tyr Ala Ala Pro Val Lys Gly Arg Phe Thr Ile Ser Arg Asp Asp Ser Lys Asn Thr Leu Tyr Leu Gln Met Asn Ser Leu Lys Thr Glu Asp Thr Ala Val Tyr 90 Tyr Cys Thr Thr <210> 609 <211> 100 <212> PRT <213> Homo sapiens <400> 609 Glu Val Gln Leu Val Glu Ser Gly Gly Ala Leu Val Lys Pro Gly Gly Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Asn Ala

Trp Met Ser Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val

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Gly Arg Ile Lys Ser Lys Thr Asp Gly Gly Thr Thr Asp Tyr Ala Ala 50 60
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Pro Val Lys Gly Arg Phe Thr Ile Ser Arg Asp Asp Ser Lys Asn Thr 65 70 75 80

Leu Tyr Leu Gln Met Asn Ser Leu Lys Thr Glu Asp Thr Ala Val Tyr 85 90 95

Tyr Cys Thr Thr 100

<210> 610

<211> 98

<212> PRT

<213> Homo sapiens

<400> 610

Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Gly
1 5 10 15

Ser Leu Arg Leu Ser Cys Pro Ala Ser Gly Phe Thr Phe Ser Asn His 20 25 30

Tyr Met Ser Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ser Tyr Ile Ser Gly Asp Ser Gly Tyr Thr Asn Tyr Ala Asp Ser Val $50 \\ 55 \\ 60$

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ala Asn Asn Ser Pro Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys $85 \hspace{1cm} 90 \hspace{1cm} 95$

Val Lys

<210> 611

<211> 98

<212> PRT

<213> Homo sapiens

<400> 611

Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Gly

1 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Asn His $20 \hspace{1cm} 25 \hspace{1cm} 30$

Tyr Thr Ser Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ser Tyr Ser Ser Gly Asn Ser Gly Tyr Thr Asn Tyr Ala Asp Ser Val 50 55 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ala Lys Asn Ser Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys 85 90 95

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Val Lys
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<210> 612

<211> 98

<212> PRT

<213> Homo sapiens

<400> 612

Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Gly
1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Asn Ser 25 30

Asp Met Asn Trp Val His Gln Ala Pro Gly Lys Gly Leu Glu Trp Val
35 40 45

Ser Gly Val Ser Trp Asn Gly Ser Arg Thr His Tyr Ala Asp Ser Val 50 60

Lys Gly Arg Phe Ile Ile Ser Arg Asp Asn Ser Arg Asn Thr Leu Tyr 65 70 75 80

Leu Gln Thr Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys
85 90 95

Val Arg

<210> 613

<211> 98

<212> PRT

<213> Homo sapiens

<400> 613

Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Gly
1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Asn Ser 20 25 30

Asp Met Asn Trp Ala Arg Lys Ala Pro Gly Lys Gly Leu Glu Trp Val
35 40 45

Ser Gly Val Ser Trp Asn Gly Ser Arg Thr His Tyr Val Asp Ser Val 50 55 60

Lys Arg Arg Phe Ile Ile Ser Arg Asp Asn Ser Arg Asn Ser Leu Tyr 65 70 75 80

Leu Gln Lys Asn Arg Arg Arg Ala Glu Asp Met Ala Val Tyr Tyr Cys 85 90 95

Val Arg

<210> 614

<211> 98

<212> PRT

<213> Homo sapiens

<400> 614

Thr Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Glu Pro Gly Gly 1 5 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Asn Ser 20 25 30

Asp Met Asn Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ser Gly Val Ser Trp Asn Gly Ser Arg Thr His Tyr Ala Asp Ser Val 50 60

Lys Gly Arg Phe Ile Ile Ser Arg Asp Asn Ser Arg Asn Phe Leu Tyr 65 70 75 90

Gln Gln Met Asn Ser Leu Arg Pro Glu Asp Met Ala Val Tyr Tyr Cys 85 90 95

Val Arg

<210> 615

<211> 97

<212> PRT

<213> Homo sapiens

<400> 615

Glu Val His Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Gly
1 5 10 15

Ala Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Asn Tyr 20 25 30

Asp Met His Trp Val Arg Gln Ala Thr Gly Lys Gly Leu Glu Trp Val 35 40 45

Ser Ala Asn Gly Thr Ala Gly Asp Thr Tyr Tyr Pro Gly Ser Val Lys 50 60

Gly Arg Phe Thr Ile Ser Arg Glu Asn Ala Lys Asn Ser Leu Tyr Leu 65 70 75 80

Gln Met Asn Ser Leu Arg Ala Gly Asp Thr Ala Val Tyr Tyr Cys Ala 85 90 95

Arg

<210> 616

<211> 97

<212> PRT

<213> Homo sapiens

<400> 616

Glu Val Gln Leu Val Glu Thr Gly Gly Gly Leu Ile Gln Pro Gly Gly
1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Val Ser Ser Asn 20 25 30

Tyr Met Ser Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val

Ser Val Ile Tyr Ser Gly Gly Ser Thr Tyr Tyr Ala Asp Ser Val Lys

- 125 -55 Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys Ala Arg <210> 617 <211> 97 <212> PRT <213> Homo sapiens <400> 617 Glu Val Gln Leu Val Gln Ser Gly Gly Gly Leu Val His Pro Gly Gly Ser Leu Arg Leu Ser Cys Ala Gly Ser Gly Phe Thr Phe Ser Ser Tyr Ala Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val Ser Ala Ile Gly Thr Gly Gly Gly Thr Tyr Tyr Ala Asp Ser Val Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ala Lys Asn Ser Leu Tyr Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Met Ala Val Tyr Tyr Cys Ala Arg <210> 618 <211> 97 <212> PRT <213> Homo sapiens <400> 618 Glu Val Gln Leu Val Gln Ser Gly Gly Gly Leu Val Gln Pro Gly Gly

Ser Leu Arg Leu Ser Cys Ala Gly Ser Gly Phe Thr Phe Ser Ser Tyr

Ala Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val

Ser Ala Ile Gly Thr Gly Gly Gly Thr Tyr Tyr Ala Asp Ser Val Lys

Gly Arg Phe Thr Ile Ser Arg Asp Asn Ala Lys Asn Ser Leu Tyr Leu

Gln Met Asn Ser Leu Arg Ala Glu Asp Met Ala Val Tyr Tyr Cys Ala 90

Arg

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<210> 619
<211> 98
<212> PRT
<213> Homo sapiens
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<400> 619

Glu Val Gln Leu Leu Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Gly
1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr
23 25 30

Ala Met Ser Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val \$35\$

Ser Ala Ile Ser Gly Ser Gly Gly Ser Thr Tyr Tyr Ala Asp Ser Val 50 55 60

Lys Gly Arg Pne Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 80

Leu Gln Met As: Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys $85 \hspace{1cm} 90 \hspace{1cm} 95$

Ala Lys

<210> 620 <211> 98 <212> PRT <213> Homo sapiens

<400> 620

Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Gly l1

Ser Leu Arg Leu Ser Cys Ser Ala Ser Gly Phe Thr Phe Ser Ser Tyr \$20\$ \$25\$ \$30

Ala Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Tyr Val 35 40 45

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Val Gln Met Ser Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys 85 90 95

Val Lys

<210> 621 <211> 98 <212> PRT <213> Homo sapiens

<400> 621 Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Gly 1 5 10 15 Ser Leu Arg Leu Ser Cys Ser Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Ala Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Tyr Val 35 40 45

Ser Ala Ile Ser Ser Asn Gly Gly Ser Thr Tyr Tyr Ala Asp Ser Val $50\,$

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Val Gln Met Ser Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys
85 90 95

Val Lys

<210> 622

<211> 98

<212> PRT

<213> Homo sapiens

<400> 622

Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Gly 1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Ala Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Tyr Val 35 40 45

Ser Ala Ile Ser Ser Asn Gly Gly Ser Thr Tyr Tyr Ala Asn Ser Val 50 55 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Gly Ser Leu Arg Ala Glu Asp Met Ala Val Tyr Tyr Cys $85 \hspace{1cm} 90 \hspace{1cm} 95$

Ala Arg

<210> 623

<211> 98

<212> PRT

<213> Homo sapiens

<400> 623

Glu Val Gln Leu Leu Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Gly
1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Ala Met Ser Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val

Ser Ala Ile Ser Gly Ser Gly Gly Ser Thr Tyr Tyr Gly Asp Ser Val $50 \hspace{1cm} 55 \hspace{1cm} 60$

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys $85 \hspace{1cm} 90 \hspace{1cm} 95$

Ala Lys

<210> 624

<211> 98

<212> PRT

<213> Homo sapiens

<400> 624

Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg
1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr
20 25 30

Ala Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val
35 40 45

Ala Val Ile Ser Tyr Asp Gly Ser Asn Lys Tyr Tyr Thr Asp Ser Val 50 55 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys $85 \hspace{1cm} 90 \hspace{1cm} 95$

Ala Arg

<210> 625

<211> 98

<212> PRT

<213> Homo sapiens

<400> 625

Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg
1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Ala Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val
35 40 45

Ala Val Ile Ser Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys
85 90 95

Ala Arg

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<210> 626
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<211> 98

<212> PRT

<213> Homo sapiens

<400> 626

Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg
1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Ala Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ala Val Ile Ser Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 30

Leu Gln Met Ser Ser Leu Arg Ala Giu Asp Thr Ala Val Tyr Tyr Cys 85 90 95

Ala Arg

<210> 627

<211> 98

<212> PRT

<213> Homo sapiens

<400> 627

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Ala Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ala Val-Ile Ser Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val50 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys $85 \hspace{1cm} 90 \hspace{1cm} 95$

Ala Arg

<210> 628

<211> 98

<212> PRT

<213> Homo sapiens

<400> 628

Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg
1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr

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20 25 30

Ala Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val
35 40 45

Ala Val Ile Ser Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys 85 90 95

Ala Arg

<210> 629

<211> 98

<212> PRT

<213> Homo sapiens

<400> 629

Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg 1 5 10

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Ala Met His Tro Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Tro Val
35 40 45

Ala Val Ile Ser Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys $85 \hspace{1cm} 90 \hspace{1cm} 95$

Ala Arg

<210> 630

<211> 98

<212> PRT

<213> Homo sapiens

<400> 630

Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg
1 5 , 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Ala Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val

Ala Val Ile Ser Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

WO 00/56772

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys $85 \hspace{1cm} 90 \hspace{1cm} 95$

Ala Arg

<210> 631

<211> 98

<212> PRT

<213> Homo sapiens

<400> 631

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Ala Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val
35 40 45

Ala Val Ile Sér Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 55 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys \$85\$ $$ 90 $$ 95

Ala Arg

<210> 632

<211> 98

<212> PRT

<213> Homo sapiens

<400> 632

Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg
1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Ala Met His Trp Val Arg Gin Ala Pro Gly Lys Gly Leu Glu Trp Val
35 40 45

Ala Val Ile Ser Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys
85 90 95

Ala Arg

<210> 633

<211> 98

<212> PRT

<213> Homo sapiens

<400> 633

Gln Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Gly
1 5 10

Ser Leu Arg Leu Ser Cys Ser Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Ala Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Tyr Val 35 40 45

Ser Ala Ile Ser Ser Asn Gly Gly Ser Thr Tyr Tyr Ala Asp Ser Val $50 \\ 55 \\ 60$

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Val Gln Met Ser Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys 85 90 95

Val Lys

<210> 634

<211> 98

<212> PRT

<213> Homo sapiens

<400> 634

Gln Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Gly
1 5 10

Ser Leu Arg Leu Se. Cys Ser Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Ala Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Tyr Val 35 40 45

Ser Ala Ile Ser Ser Asn Gly Gly Ser Thr Tyr Tyr Ala Asp Ser Val 50 55 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys
85 90 95

Ala Arg ·

<210> 635

<211> 98

<212> PRT

<213> Homo sapiens

<400> 635

Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg 1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Ala Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ala Val Ile Ser Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 60

Lys Gly Arg Phe Ala Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 . 70 75

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys 85 90 95

Ala Arg

<21.0> 636

<211> 98

<212> PRT

<213> Homo sapiens

<400> 636

Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg 1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Ala Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ala Val Ile Ser Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 55 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 30

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys $85 \hspace{1cm} 90 \hspace{1cm} 95$

Ala Arg

<210> 637

<211> 98

<212> PRT

<213> Homo sapiens

<400> 637

Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg 1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Ala Met His Trp Val Arg Gin Ala Pro Gly Lys Gly Leu Glu Trp Val
35 40 45

Ala Val Ile Ser Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys

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85 90 95

Ala Lys

<210> 638

<211> 97

<212> PRT

<213> Homo sapiens

<400> 638

Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Gly 1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Asp Met His Trp Val Arg Gln Ala Thr Gly Lys Gly Leu Glu Trp Val 35 40 45

Ser Ala Ile Gly Thr Ala Gly Asp Thr Tyr Tyr Pro Gly Ser Val Lys 50 60

Gly Arg Phe Thr Ile Ser Arg Glu Asn Ala Lys Asn Ser Leu Tyr Leu 65 70 75 80

Gln Met Asn Ser Leu Arg Ala Gly Asp Thr Ala Val Tyr Tyr Cys Ala 85 90 95

Arg

<210> 639

<211> 98

<212> PRT

<213> Homo sapiens

<400> 639

Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Prc Gly Gly
1 5 10

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Glu Met Asn Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val
35 40 45

Ser Tyr Ile Ser Ser Ser Gly Ser Thr Ile Tyr Tyr Ala Asp Ser Val 50 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ala Lys Asn Ser Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys $85 \hspace{1cm} 90 \hspace{1cm} 95$

Ala Arg

<210> 640

<211> 98

<212> PRT

<213> Homo sapiens

<400> 640

Gin Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg
1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val
35 40 45

Ala Val Ile Ser Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Leu Arg Ala Arg Leu Cys Ile Thr Val 85 90 95

Arg Glu

<210> 641

<211> 98

<212> PRT

<213> Homo sapiens

<400> 641

Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg
1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ala Val Ile Ser Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys 85 90 95

Ala Arg

<210> 642

<211> 98

<212> PRT

<213> Homo sapiens

<400> 642

Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45 Ala Val Ile Ser Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Rsn Ser Lys Asn Thr Leu Tyr 65 , 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys $85 \hspace{1cm} 90 \hspace{1cm} 95$

Ala Arg

<210> 643

<211> 98

<212> PRT

<213> Homo sapiens

. <400> 643

Gin Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg 1 5 10

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val
35 40 45

Ala Val Ile Ser Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 55 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys $85 \hspace{1cm} 90 \hspace{1cm} 95$

Ala Arg

<210> 644

<211> 98

<212> PRT

<213> Homo sapiens

<400> 644

Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg 1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ala Val Ile Ser Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys $85 \hspace{1.5cm} 90 \hspace{1.5cm} 95$

Ala Arg

<210> 645

<211> 98

<212> PRT

<213> Homo sapiens

<400> 645

Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg 1 $$. 5 $$ 10

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr
20 25 30

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val
35 40 45

Ala Val Ile Ser Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 55 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Arg Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys 85 90 95

Ala Arg

<210> 646

<211> 98

<212> PRT

<213> Homo sapiens

<400> 646

Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg 1 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val
35 40 45

Ala Val Ile Ser Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys
85 90 95

Ala Arg

<210> 647

<211> 98

<212> PRT

<213> Homo sapiens

<400> 647

- 138 -

Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg 1 5 10

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr $20 \hspace{1cm} 25 \hspace{1cm} 30$

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val
35 40 45

Ala Val Ile Trp Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys $85 \hspace{1cm} 90 \hspace{1cm} 95$

Ala Arg

<210> 648

<211> 98

<212> PRT

<213> Homo sapiens

<400> 648

Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg 1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 . 30

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ala Val Ile Ser Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Gly Thr Ala Val Tyr Tyr Cys . $85 \hspace{1cm} 90 \hspace{1cm} 95$

Ala Arg

<210> 649

<211> 98

<212> PRT

<213> Homo sapiens

<400> 649

Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Gly
1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 \cdot 25 30

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ala Phe Ile Arg Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val

- 139 -

50 55 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys \$85\$ \$90 \$95

Ala Lys

<210> 650

<211> 98

<212> PRT

<213> Homo sapiens

<400> 650

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ala Val Ile Ser Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys $85 \hspace{1cm} 90 \hspace{1cm} 95$

Ala Lys

<210> 651

<211> 98

<212> PRT

<213> Homo sapiens

<400> 651

Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ala Val Ile Trp Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys 85 90 95

Ala Arg

- 140 -

<210> 652 <211> 98 <212> PRT <213> Homo sapiens <400> 652 Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg 10 Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val Ala Val Ile Trp Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr-Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys Ala Lys <210> 653 <211> 95 <212> PRT <213> Homo sapiens <400> 653 Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val Ala Val Ile Ser Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 55 Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Arg Lys <210> 654

<211> 98 <212> PRT <213> Homo sapiens <400> 654 Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr

- 141 -

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ala Val Ile Ser Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys 85 90 95

Ala Arg

<210> 655

<211> 98

<212> PRT

<213> Homo sapiens

<400> 655

Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg
1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ala Val Ile Trp Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Ala 50 55 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Thr Asn Thr Leu Phe 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys $85 \hspace{1cm} 90 \hspace{1cm} 95$

Ala Arg

<210> 656

<211> 98

<212> PRT

<213> Homo sapiens

<400> 656

Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Gly
1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Ser Met Asn Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val
35 40 45

Ser Tyr Ile Ser Ser Ser Ser Ser Thr Ile Tyr Tyr Ala Asp Ser Val

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ala Lys Asn Ser Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Asp Glu Asp Thr Ala Val Tyr Tyr Cys $85 \hspace{1cm} 95$

Ala Arg

<210> 657

<211> 98

<212> PRT

<213> Homo sapiens

<400> 657

Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Lys Pro Gly Gly
1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Ser Met Asn Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ser Ser Ile Ser Ser Ser Ser Ser Tyr Île Tyr Tyr Ala Asp Ser Val $50 \hspace{1.5cm} 55 \hspace{1.5cm} 60$

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ala Lys Asn Ser Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys $85 \hspace{1cm} 90 \hspace{1cm} 95$

Ala Arg

<210> 658

<211> 97

<212> PRT

<213> Homo sapiens

<400> 658

Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Lys Pro Gly Gly 1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Ser Met Asn Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ser Ser Ile Ser Ser Ser Ser Tyr Ile Tyr Tyr Ala Asp Ser Val Lys 50 55 60

Gly Arg Phe Thr Ile Ser Arg Asp Asn Ala Lys Asn Ser Leu Tyr Leu 65 70 75 80

Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys Ala 85 90 95

Arg

<210> 659

<211> 98

<212> PRT

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<213> Homo sapiens
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<400> 659

Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Lys Pro Gly Gly

1 5 • 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Ser Met Asn Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ser Ser Ile Ser Ser Ser Ser Tyr Ile Tyr Tyr Ala Asp Ser Val
50 55 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ala Lys Asn Ser Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys $85 \hspace{1cm} 90 \hspace{1cm} 95$

Ala Arg

<210> 660

<211> 98

<212> PRT

<213> Homo sapiens.

<400> 660

Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Gly
1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Ser Met Asn Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ser Tyr Ile Ser Ser Ser Ser Thr Ile Tyr Tyr Ala Asp Ser Val 50 55 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ala Lys Asn Ser Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys $85 \hspace{1cm} 90 \hspace{1cm} 95$

Ala Arg

<210> 661

<211> 97

<212> PRT

<213> Homo sapiens

<400> 661

Glu Asp Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Gly
1 5 10

Ser Leu Arg Pro Ser Cys Ala Ala Ser Gly Phe Ala Phe Ser Ser Tyr

Val Leu His Trp Val Arg Arg Ala Pro Gly Lys Gly Pro Glu Trp Val

- 144 -

35 40 . 45

Ser Ala Ile Gly Thr Gly Gly Asp Thr Tyr Tyr Ala Asp Ser Val Met 50 . 60

Gly Arg Phe Thr Ile Ser Arg Asp Asn Ala Lys Lys Ser Leu Tyr Leu 65 70 75 30

Gln Met Asn Ser Leu Ile Ala Glu Asp Met Ala Val Tyr Tyr Cys Ala 85 90

Arg

<210> 662

<211> 98

<212> PRT

<213> Homo sapiens

<400> 662

Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Gly
1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Trp Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Val Trp Val 35 40 45

Ser Arg Ile Asn Ser Asp Gly Ser Ser Thr Ser Tyr Ala Asp Ser Val 50 55 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ala Lys Asn Thr Leu Tyr 65 . 70 . 75

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys 85 90 95

Ala Arg

<210> 663

<211> 98

<212> PRT

<213> Homo sapiens

<400> 663

Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Gly
1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Trp Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Val Trp Val 35 40 45

Ser Arg Ile Asn Ser Asp Gly Ser Ser Thr Ser Tyr Ala Asp Ser Val

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ala Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys $85 \hspace{1cm} 90 \hspace{1cm} 95$

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Ala Arg
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<210> 664

<211> 98

<212> PRT

<213> Homo sapiens

<400> 664

Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Gly 1 $$ 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Trp Met Ser Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ala Lys Asn Ser Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys 85 90 95

Ala Arg

<210> 665

<211> 98

<212> PRT

<213> Homo sapiens

<400> 665

Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Gly
1 5 10

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Trp Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Val Trp Val 35 40 45

Ser Arg Ile Asn Ser Asp Gly Ser Ser Thr Ser Tyr Ala Asp Ser Met 50 60

Lys Gly Gln Phe Thr Ile Ser Arg Asp Asn Ala Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Met Ala Val Tyr Tyr Cys 85 90 95

Thr Arg

<210> 666

<211> 98

<212> PRT

<213> Homo sapiens

- 146 -

<400> 666

Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Gly 1 5 10 15

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Trp Met Ser Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 35 40 45

Ala Asn Ile Lys Gln Asp Gly Ser Glu Lys Tyr Tyr Val Asp Ser Val 50 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ala Lys Asn Ser Leu Tyr 65 70 75 80

Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys 85 90 95

Ala Arg

<210> 667

<211> 98

<212> PRT

<213> Homo sapiens

<400> 667

Gln Val Gln Leu Val Gln Ser Gly Gly Gly Val Val Gln Pro Gly Arg 1 5 10

Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr 20 25 30

Gly Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val
35 40 45

Ala Phe Ile Arg Tyr Asp Gly Ser Asn Lys Tyr Tyr Ala Asp Ser Val 50 55 60

Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Tyr 65 70 75 80

Leu Gln Met Lys Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr Tyr Cys 85 90 95

Thr Thr

<210> 668

<211> 98

<212> PRT <213> Homo sapiens

<400> 668

Gln Ser Val Leu Thr Gln Pro Pro Ser Val Ser Ala Ala Pro Gly Gln $1 \hspace{1cm} 5 \hspace{1cm} 10 \hspace{1cm} 15$

Lys Val Thr Ile Ser Cys Ser Gly Ser Ser Ser Asn Ile Gly Asn Asn

Tyr Val Ser Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu 35 40 45

- 147 -

Ile Tyr Asp Asn Asn Lys Arg Pro Ser Gly Ile Pro Asp Arg Phe Ser 50 60

Gly Ser Lys Ser Gly Thr Ser Ala Thr Leu Gly Ile Thr Gly Leu Gln 65 70 75 80

Thr Gly Asp Glu Ala Asp Tyr Tyr Cys Gly Thr Trp Asp Ser Ser Leu 85 90 95

Ser Ala

<210> 669

<211> 98

<212> PRT

<213> Homo sapiens

<400> 669

Gln Ser Val Leu Thr Gln Pro Pro Ser Val Ser Ala Ala Pro Gly Gln
1 5 10 15

Lys Val Thr Ile Ser Cys Ser Gly Ser Ser Ser Asp Met Gly Asn Tyr 20 25 30

Ala Val Ser Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu $35 \hspace{1.5cm} 40 \hspace{1.5cm} 45$

Ile Tyr Glu Asn Asn Lys Arg Pro Ser Gly Ile Pro Asp Arg Phe Ser
50 55 60

Gly Ser Lys Ser Gly Thr Ser Ala Thr Leu Gly Ile Thr Gly Leu Trp 65 75 80

Pro Glu Asp Glu Ala Asp Tyr Tyr Cys Leu Ala Trp Asp Thr Ser Pro
85 90 95

Arg Ala

<210> 670

<211> 98

<212> PRT

<213> Homo sapiens

<400> 670

Gln Ser Val Leu Thr Gln Pro Pro Ser Ala Ser Gly Thr Pro Gly Gln 1 5 10 15

Arg Val Thr Ile Ser Cys Ser Gly Ser Ser Ser Asn Ile Gly Ser Asn 20 25 30

Thr Val Asn Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu
35 40

Ile Tyr Ser Asn Asn Gln Arg Pro Ser Gly Val Pro Asp Arg Phe Ser

Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Ser Gly Leu Gln 65 70 75 80

Ser Glu Asp Glu Ala Asp Tyr Tyr Cys Ala Ala. Trp Asp Asp Ser Leu 85 90 95

Asn Gly

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<210> 671
<211> 98
<212> PRT
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<213> Homo sapiens

<400> 671

Gin Ser Val Leu Thr Gln Pro Pro Ser Ala Ser Gly Thr Pro Gly Gln 1 5 10 15

Arg Val Thr Ile Ser Cys Ser Gly Ser Ser Ser Asn Ile Gly Ser Asn 20 25 30

Tyr Val Tyr Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu 35 40 45

Ile Tyr Arg Asn Asn Gln Arg Pro Ser Gly Val Pro Asp Arg Phe Ser 50 60

Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Ser Gly Leu Arg 65 70 75 80

Ser Glu Asp Glu Ala Asp Tyr Tyr Cys Ala Ala Trp Asp Asp Ser Leu $85 \hspace{1cm} 90 \hspace{1cm} 95$

Ser Gly

<210> 672

<211> 98 <212> PRT

<213> Homo sapiens

<400> 672

Gln Ser Val Leu Thr Gln Pro Pro Ser Val Ser Glu Ala Pro Arg Gln $1 \hspace{1cm} 5 \hspace{1cm} 10 \hspace{1cm} 15$

Arg Val Thr Ile Ser Cys Ser Gly Ser Ser Ser Asn Ile Gly Asn Asn 20 25 30

Ala Val Asn Trp Tyr Gln Gln Leu Pro Gly Lys Ala Pro Lys Leu Leu $35 \hspace{1cm} 40 \hspace{1cm} 45$

Ile Tyr Tyr Asp Asp Leu Leu Pro Ser Gly Val Ser Asp Arg Phe Ser 50 60

Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Ser Gly Leu Gln 65 70 75 . 80

Ser Glu Asp Glu Ala Asp Tyr Tyr Cys Ala Ala Trp Asp Asp Ser Leu 85 90 95

Asn Gly

<210> 673

<211> 99

<212> PRT

<213> Homo sapiens

<400> 673

Gln Ser Val Leu Thr Gln Pro Pro Ser Val Ser Gly Ala Pro Gly Gln

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10 1.5 Arg Val Thr Ile Ser Cys Thr Gly Ser Ser Ser Asn Ile Gly Ala Gly Tyr Val Val His Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu Ile Tyr Gly Asn Ser Asn Arg Pro Ser Gly Val Pro Asp Gln Phe Ser Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Thr Gly Leu Gln Ser Glu Asp Glu Ala Asp Tyr Tyr Cys Lys Ala Trp Asp Asn Ser Leu Asn Ala <210> 674 <211> 99 <212> PRT <213> Homo sapiens <400> 674 Gln Ser Val Val Thr Gln Pro Pro Ser Val Ser Gly Ala Pro Gly Gln Arg Val Thr Ile Ser Cys Thr Gly Ser Ser Ser Asn Ile Gly Ala Gly Tyr Asp Val His Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu 40 Leu Ile Tyr Gly Asn Ser Asn Arg Pro Ser Gly Val Pro Asp Arg Phe Ser Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Thr Gly Leu Gln Ala Glu Asp Glu Ala Asp Tyr Tyr Cys Gln Ser Tyr Asp Ser Ser Leu Ser Gly <210> 675 <211> 98 <212> PRT <213> Homo sapiens <400> 675 Ser Tyr Val Leu Thr Gln Pro Pro Ser Val Ser Gly Thr Pro Gly Gln 10

Arg Val Thr Ile Ser Cys Ser Gly Gly Arg Ser Asn Ile Gly Ser Asn 20 25 30

Thr Val Lys Trp Tyr Gln Gln Leu Pro Gly Thr Ala Pro Lys Leu Leu 35 40 45

Ile Tyr Gly Asn Asp Gln Arg Pro Ser Gly Val Pro Asp Arg Phe Ser 50 60

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- 150 -

Gly Ser Lys Ser Gly Thr Ser Ala Ser Leu Ala Ile Thr Gly Val Gln 65 70 . 75 80

Ala Glu Asp Glu Ala Asp Tyr Tyr Cys Gln Ser Tyr Asp Ser Ser Leu $85 \ \cdot \ 90 \ 95$

Arg Gly

INTERNATIONAL SEARCH REPORT

International Application No PCT/US 00/07946

PCT/US 00/07946 A. CLASSIFICATION OF SUBJECT MATTER IPC 7 C07K16/24 C12N C12N5/10 C07K16/00 C12N15/13 C12N15/63 A61K39/395 C12P21/08 A61P43/00 G01N33/577 According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) IPC 7 C07K Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) STRAND, MEDLINE, BIOSIS, WPI Data, EPO-Internal, PAJ C. DOCUMENTS CONSIDERED TO BE RELEVANT Relevant to claim No. Category Citation of document, with indication, where appropriate, of the relevant passages Α EP 0 659 766 A (SCHERING PLOUGH CORP) 28 June 1995 (1995-06-28) page 2, line 47-54 page 3, line 34 -page 4, line 33 claims Α WO 94 04679 A (GENENTECH INC) 3 March 1994 (1994-03-03) page 5, line 1 -page 7, line 3 example 3 claims Further documents are listed in the continuation of box C. Patent family members are listed in annex. X X Special categories of cited documents: "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the "A" document defining the general state of the art which is not considered to be of particular relevance invention *E" earlier document but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone filing date document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another "Y" document of particular relevance; the claimed invention citation or other special reason (as specified) cannot be considered to involve an inventive step when the "O" document referring to an oral disclosure, use, exhibition or document is combined with one or more other such docu-ments, such combination being obvious to a person skilled other means "P" document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 8 August 2000 23/08/2000 Authorized officer Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl,

Fax: (+31-70) 340-3016

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Covone, M

INTERNATIONAL SEARCH REPORT

International Application No
PCT/US 00/07946

		PC1/03 00	
C.(Continua	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	·	
Category °	Citation of document, with indication,where appropriate, of the relevant passages		Relevant to claim No.
A	DATABASE BIOSIS 'Online! BIOSCIENCES INFORMATION SERVICE, PHILADELPHIA, PA, US; 1997 CARTER R W ET AL: "Production and characterization of monoclonal antibodies to human interleukin-12." Database accession no. PREV199799787174 XP002144460 abstract & HYBRIDOMA, vol. 16, no. 4, 1997, pages 363-369, ISSN: 0272-457X		
Α	PINI A ET AL: "Hierarchical affinity maturation of a phage library derived antibody for the selective removal of cytomegalovirus from plasma" JOURNAL OF IMMUNOLOGICAL METHODS, NL, ELSEVIER SCIENCE PUBLISHERS B.V., AMSTERDAM, vol. 206, no. 1-2, 7 August 1997 (1997-08-07), pages 171-182, XP004093130 ISSN: 0022-1759 the whole document		
Α	IRVING R A ET AL: "Affinity maturation of recombinant antibodies using E.colimutator cells" IMMUNOTECHNOLOGY,NL,ELSEVIER SCIENCE PUBLISHERS BV, vol. 2, no. 2, 1 June 1996 (1996-06-01), pages 127-143, XP004052677 ISSN: 1380-2933 abstract figure 1		
A	WO 95 24918 A (GENETICS INST) 21 September 1995 (1995-09-21) page 3, line 15 -page 4, line 14		

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FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

Continuation of Box I.1

Although claims 94,95,142 (completely) are directed to a method of treatment of the human/animal body, and claim 139 (partially) is directed to a diagnostic method practised on the human/animal body, the search has been carried out and based on the alleged effects of the compound/composition.

Continuation of Box I.2

In view of the large number and also the wording of the claims presently on file, which render it difficult, if not impossible, to determine the matter for which protection is sought, the present application fails to comply with the clarity and conciseness requirements of Article 6 PCT (see also Rule 6.1(a) PCT) to such an extent that a meaningful search is impossible. Moreover support within the meaning of Article 6 PCT and/or disclosure within the meaning of Article 5 PCT is to be found for only a very small proportion of the compounds and methods claimed. In the present case, the claims also so lack support, and the application so lacks disclosure, that a meaningful search over the whole of the claimed scope is impossible.

Consequently, the search has been carried out for those parts of the application which do appear to be clear, concise and supported by the experimental data, namely those parts related to human anti-IL-12 antibodies as disclosed at page 140 line 20 - page 141 line 10 (antibody Y61) and at page 143 line 12-line 32 (antibody J695) and in claims 23-26 and 41-44, including nucleic acid coding for said antibodies, host cell to express said nucleic acid and production of said protein. Thus claims 23-31, 35-45, 47-49, 53, 55-58, 60-62, 64, 66-69, 71-73 have been fully searched. The remaining claims have been partially searched, in the light of the supported subject matter.

The applicant's attention is drawn to the fact that claims, or parts of claims, relating to inventions in respect of which no international search report has been established need not be the subject of an international preliminary examination (Rule 66.1(e) PCT). The applicant is advised that the EPO policy when acting as an International Preliminary Examining Authority is normally not to carry out a preliminary examination on matter which has not been searched. This is the case irrespective of whether or not the claims are amended following receipt of the search report or during any Chapter II procedure.

INTERNATIONAL SEARCH REPORT

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